FIGURE 1

GCCCACGCGTCCGATGGCGTTCACGTTCGCGGCCTTCTGCTACATGCTGGCGCTGCTGCT CACTGCCGCGCTCATCTTCTCCGCCATTTGGCACATTATAGCATTTGATGAGCTGAAGAC TGATTACAAGAATCCTATAGACCAGTGTAATACCCTGAATCCCCTTGTACTCCCAGAGTA CCTCATCCACGCTTTCTTCTGTGTCATGTTTCTTTGTGCAGCAGAGTGGCTTACACTGGG TCTCAATATGCCCCTCTTGGCATATCATATTTGGAGGTATATGAGTAGACCAGTGATGAG TGGCCCAGGACTCTATGACCCTACAACCATCATGAATGCAGATATTCTAGCATATTGTCA GAAGGAAGGATGGTGCAAATTAGCTTTTTATCTTCTAGCATTTTTTTACTACCTATATGG CATGATCTATGTTTTGGTGAGCTCTTAGAACAACACACAGAAGAATTGGTCCAGTTAAGT GCATGCAAAAAGCCACCAAATGAAGGGATTCTATCCAGCAAGATCCTGTCCAAGAGTAGC TTTTGCTTGTGGAAAGACTGTTTTCATATGTTATACTCAGATAAAGATTTTAAATGGTAT TACGTATAAATTAATATAAAATGATTACCTCTGGTGTTGACAGGTTTGAACTTGCACTTC GAAGCTTTTGTTTATAGGAACTTGTAGGGCTCATTTTGGTTTCATTGAAACAGTATCTAA TTATAAATTAGCTGTAGATATCAGGTGCTTCTGATGAAGTGAAAATGTATATCTGACTAG TGGGAAACTTCATGGGTTTCCTCATCTGTCATGTCGATGATTATATATGGATACATTTAC AAAAATAAAAAGCGGGAATTTTCCCTTCGCTTGAATATTATCCCTGTATATTTGCATGAAT GAGAGATTTCCCATATTTCCATCAGAGTAATAAATATACTTGCTTTAATTCTTAAGCATA AGTAAACATGATATAAAAATATATGCTGAATTACTTGTGAAGAATGCATTTAAAGCTATT TTAAATGTGTTTTTATTTGTAAGACATTACTTATTAAGAAATTGGTTATTATGCTTACTG TTCTAATCTGGTGGTAAAGGTATTCTTAAGAATTTGCAGGTACTACAGATTTTCAAAACT GAATGAGAGAAAATTGTATAACCATCCTGCTGTTCCTTTAGTGCAATACAATAAAACTCT GAAATTAAGACTC

FIGURE 3

TTCGCTGGATTCAGGGTCTTGGGCATCAGAGGTGAGAGGGTGGGAAGGTCCGCCGCGATG GGGAAGCCCTGGCTGCGCTGCAGCTGCTGCTGCTGGGCGCGCGTCGTGGGCGCGC GCGGGCGCCCCGCGCTGCACCTACACCTTCGTGCTGCCCCCGCAGAAGTTCACGGGCGCT GTGTGCTGGAGCGCCCCGCATCCACGCGGGCGACGCCCGAGGCCGCCAACGCCAGCGAG CTGGCGCGCTGCGCATGCGCGTCGGCCGCCACGAGGAGCTGTTACGCGAGCTGCAGAGG CTGGCGGCCGACGGCGCGTGGCCGAGGTGCGCGCGCTGCGCAAGGAGAGCCGC GGCCTGAGCGCGCCCTGGGCCAGTTGCGCGCGCAGCTGCAGCACGAGGCGGGCCCGGG CGCGTGCTCAACGCGTCCGCCGAGGCTCAGCGCGCAGCCGGCTCCACCAGCTGGAC GAGCGCCTGTGCCCGGGAGGCGGGCGGGCAGCAGCAGGTCCTGCCGCCACCCCACTG GTGCCTGTGGTTCCGGTCCGTCTTGTGGGTAGCACCAGTGACACCAGTAGGATGCTGGAC CCAGCCCCAGAGCCCCAGAGACCCAGAGCCAGGCCCATGGCTTCTCCC ATGCCTGCAGGTCACCCTGCGGTCCCCACCAAGCCTGTGGGCCCGTGGCAGGATTGTGCA GAGGCCCGCCAGGCAGGCCATGAACAGAGTGGAGTGTATGAACTGCGAGTGGGCCGTCAC GTAGTGTCAGTATGGTGTGAGCAGCAACTGGAGGGTGGAGGCTGGACTGTGATCCAGCGG AGGCAAGATGGTTCAGTCAACTTCTTCACTACCTGGCAGCACTATAAGGCGGGCTTTTGGC CGGCCAGACGGAGAATACTGGCTGGGCCTTGAACCCGTGTATCAGCTGACCAGCCGTGGG GACCATGAGCTGCTGGTTCTCCTGGAGGACTGGGGGGGCCGTGGAGCACGTGCCCACTAT GATGGCTTCTCCCTGGAACCCGAGAGCGACCACTACCGCCTGCGGCTTGGCCAGTACCAT GGTGATGCTGGAGACTCTCTTTCCTGGCACAATGACAAGCCCTTCAGCACCGTGGATAGG GACCGAGACTCCTATTCTGGTAACTGTGCCCTGTACCAGCGGGGAGGCTGGTGGTACCAT GCCTGTGCCCACTCCAACCTCAACGGTGTGTGGCACCACGGCGGCCACTACCGAAGCCGC ${\tt TACCAGGATGGTGTCTACTGGGCTGAGTTTCGTGGTGGGGGCATATTCTCTCAGGAAGGCC}$ GCCATGCTCATTCGGCCCCTGAAGCTGTGACTCTGTGTTCCTCTGTCCCCTAGGCCCTAG AGGACATTGGTCAGCAGGAGCCCAAGTTGTTCTGGCCACACCTTCTTTGTGGCTCAGTGC CAATGTGTCCCACAGAACTTCCCACTGTGGATCTGTGACCCTGGGCGCTGAAATGGGAC CGGTTTGAGCTCATATCTTATAATAACACAAAGTAGCCAC

FIGURE 4

MGKPWLRALOLLLLLGASWARAGAPRCTYTFVLPPQKFTGAVCWSGPASTRATPEAANAS
ELAALRMRVGRHEELLRELQRLAAADGAVAGEVRALRKESRGLSARLGQLRAQLQHEAGP
GAGFGADLGAEPAAALALLGERVLNASAEAQRAAARFHQLDVKFRELAQLVTQGSSLIAR
LERLCPGGAGGQQOVLPPPPLVPVVPVRLVGSTSDTSRMLDPAPEPQRDQTQRQQEPMAS
PMPAGHPAVPTKPUGPWQDCAEARQAGHEGSGVYELEVGRHVVSVWGEQOLEGGGWTVIQ
RRQDGSVNFFTTWQHYKAGFGRPDGEYWLGLEPVYQLTSRGDHELLVLLEDWGGRGARAH
YDGFSLEPESDHYRLRLGQYHGDAGDSLSWHNDKPFSTVDRDRDSYSGNCALYQRGGWWY
HACAHSNLNGVWHHGGHYRSRYODGVYWAEFFGGAYSLRKAAMLIRPLKL

Signal peptide: Amino acids 1-20

N-glycosylation sites: Amino acids 58-62;145-149

 $\mathtt{cAMP-}$ and $\mathtt{cGMP-} \mathtt{dependent}$ protein kinase phosphorylation site:

Amino acids 97-101

Tyrosine kinase phosphorylation site: Amino acids 441-448

N-myristoylation sites: Amino acids

16-22;23-29;87-93;108-114;121-127;125-131;129-135;187-193;29 3-299;353-359;378-384;445-451;453-459

Cell attachment sequence: Amino acids 340-343

Fibrinogen beta and gamma chains C-terminal domain signature:

Amino acids 418-431

FIGURE 5

CCCACGCGTCCGGCGCGTGGCCTCGCGTCCATCTTTGCCGTTCTCTCGGACCTGTCACA AAGGAGTCGCGCCGCCGCCCCCCCCCCCCCCCCCCGGGGGCCCGGAGGTAGAGAAAGT GGACTGGGGCCGGGGTAGGCTCTGGAAAGGGCCCGGGAGAGAGGTGGCGTTGGTCAGAAC CTGAGAAACAGCCGAGAGGTTTTCCACCGAGGCCCGCGCTTGAGGGATCTGAAGAGGTTC CTAGAAGAGGGTGTTCCCTCTTTCGGGGGTCCTCACCAGAAGAGGTTCTTGGGGGTCGCC CTTCTGAGGAGGCTGCGGCTAACAGGGCCCAGAACTGCCATTGGATGTCCAGAATCCCCT AAATAGGATGCAAATTCCTCAACTCCAGGTTATGAAAACAGTACTTGGAAAACTGAAAAC TACCTAAATGATCGTCTTTGGTTGGGCCGTGTTCTTAGCGAGCAGAAGCCTTGGCCAGGG TCTGTTGTTGACTCTCGAAGAGCACATAGCCCACTTCCTAGGGACTGGAGGTGCCGCTAC TACCATGGGTAATTCCTGTATCTGCCGAGATGACAGTGGAACAGATGACAGTGTTGACAC CCAACAGCAACAGGCCGAGAACAGTGCAGTACCCACTGCTGACACAAGGAGCCAACCACG AAATGTGGATGGGCTAGTGTTGGACACACTGGCAGTAATACGGACTCTTGTAGATAAGTA AGTATCTGACTCACGGTCACCTCCAGTGGAATGAAAAGTGTTCTGCCCGGAACCATGACT TTAGGACTCCTTCAGTTCCTTTAGGACATACTCGCCAAGCCTTGTGCTCACAGGGCAAAG GAGAATATTTTAATGCTCCGCTGATGGCAGAGTAAATGATAAGATTTGATGTTTTTGCTT GCTGTCATCTACTTTGTCTGGAAATGTCTAAATGTTTCTGTAGCAGAAAACACGATAAAG CTATGATCTTTATTAGAG

FIGURE 6

MIVFGWAVFLASRSLGQGLLLTLEEHIAHFLGTGGAATTMGNSCICRDDSGTDDSVDTQQ OOAENSAVPTADTRSOPRDPVRPPRRGRGPHEPRRKKONVDGLVLDTLAVIRTLVDKO

Signal peptide: amino acids 1-16

Casein kinase II phosphorylation site: amino acids 22-26, 50-54, 113-117

N-myristoylation site: amino acids 18-24, 32-38, 34-40, 35-41, 51-57

FIGURE 7

CGGACGCGTGGGGGAAACCCTTCCGAGAAAACAGCAACAAGCTGAGCTGCTGTGACAGAG GGGAACAAGATGGCGGCGCCGAAGGGGAGCCTCTGGGTGAGGACCCAACTGGGGCTCCCG CCGCTGCTGCTGACCATGGCCTTGGCCGGAGGTTCGGGGACCGCTTCGGCTGAAGCA TTTGACTCGGTCTTGGGTGATACGGCGTCTTGCCACCGGGCCTGTCAGTTGACCTACCCC TTGCACACCTACCCTAAGGAAGAGGAGTTGTACGCATGTCAGAGAGGGTTGCAGGCTGTTT TCAATTTGTCAGTTTGTGGATGATGGAATTGACTTAAATCGAACTAAATTGGAATGTGAA TCTGCATGTACAGAAGCATATTCCCAATCTGATGAGCAATATGCTTGCCATCTTGGTTGC CAGAATCAGCTGCCATTCGCTGAACTGAGACAAGAACAACTTATGTCCCTGATGCCAAAA ATGCACCTACTCTTTCCTCTAACTCTGGTGAGGTCATTCTGGAGTGACATGATGGACTCC GCACAGAGCTTCATAACCTCTTCATGGACTTTTTATCTTCAAGCCGATGACGGAAAAATA GTTATATTCCAGTCTAAGCCAGAAATCCAGTACGCACCACATTTGGAGCAGGAGCCTACA AATTTGAGAGAATCATCTCTAAGCAAAATGTCCTATCTGCAAATGAGAAATTCACAAGCG TCTGGGTGGATTTTAACTACAACTCTTGTCCTCTCGGTGATGGTATTGCTTTGGATTTGT TGTGCAACTGTTGCTACAGCTGTGGAGCAGTATGTTCCCTCTGAGAAGCTGAGTATCTAT GGTGACTTGGAGTTTATGAATGAACAAAAGCTAAACAGATATCCAGCTTCTTCTCTTGTG $\mathtt{CTTGCTCATTCTGAAAT}$ AATTCCACTCCTCATAGAGCTTTTAAAATGGTTTCATTGGATATAGGCCTTAAGAAATCA CTATAAAATGCAAATAAAGTTACTCAAATCTGTG

FIGURE 8

MAAPKGSLWVRTQLGLPPLLLLTMALAGGSGTASAEAFDSVLGDTASCHRACQLTYPLHT YPKEEELYACQRGCRLFSICQFVDDGIDLNRTKLECESACTEAYSQSDEQYACHLGCQND LPFAELROEQLMSLMPKMHLLFPLTLVRSFWSDMMDSAQSFITSSWTFYLQADDGKIVIF QSKPEIQYAPHLEQEPTNLRESSLSKMSYLQMRNSQAHRNFLEDGESDGFLRCLSLNSGW ILTTTLVLSVMVLLWICCATVATAVEQYVPSEKLSIYGDLEFMNEQKLNRYPASSLVVVR SKTEDHEEAGPLPTKVNLAHSEI

Important features: Signal peptide: amino acids 1-31

Transmembrane domain: amino acids 241-260

N-glycosylation site: amino acids 90-93

FIGURE 9

FIGURE 10

MKGSIFTLFLFSVLFAISEVRSKESVRLCGLEYIRTVIYICASSRWRRHLEGIPQAQQAE TGNSFQLPHKREFSEENPAQNLPKVDASGEDRLWGGQMPTEELWKSKKHSVMSRQDLQTL CCTDGCSWTDLSALC

Important features: Signal sequence: amino acids 1-18

 $\mathtt{cAMP-}$ and $\mathtt{cGMP-} \mathtt{dependent}$ protein kinase phosphorylation site:

amino acids 107-111

N-myristoylation sites: amino acids 3-9,52-58,96-102,125-131

Insulin family signature: amino acids 121-136

Insulin family proteins: amino acids 28-46

FIGURE 11

CCCACGCGTCCGGACAAACTGGAGGTGAAAGGAGCTGGTACTGTCCACTGTGCTGTCGGT GCTGAACCTGAGACGCGAGCGGACCAGTTGCTCCAGCACCTGAAGGCAACGCCCTCTTGC ACCCTCTGTGCCCTGTGGGACCCGCTTCACCAACAGGACCCATATCAACTTGACAAAGGA GTGTGGTATCGGACGTGGGAGAGAGTCCTCTGTTTGCCACCTGGGCGCTCATTCAGGCGT CTTTTGAGAGTTTTTTTTTTTTTTTGTGATTTATATTTCCACAGCGTTTAGGAATCTTTCT GGGGGACTTTTGTGACTGTTAAAATAAGGTGAAAAGCAATAAGGATG GAGGATATGGCCTTAAATCCTATCAGCCTCTAATGAGATTGCGACATAAGCAGGAAAAAA ATCAAGAAAGTTCAAGAGTCAAAGGATTTATGATTCAGGATGGCCCTTTTGGATCTTGTG AAAATAAGTACTGTGGTTTGGGAAGACACTGTGTTACCAGCAGAGAGACAGGGCAAGCAG AATTCTATGAAAACCACTGTGAAGTGCACAGAGCTGCTTGCCTGAAAAAAACAAAAGATTA CCATTGTTCACAATGAAGACTGCTTCTTTAAAGGAGATAAGTGCAAGACTACTGAATACA GCAAGATGAAAAATATGCTATTAGATTTACAAAATCAAAAATATATTATGCAAGAAAATG AAAATCCTAATGGCGACGACATATCTCGGAAGAAGCTATTGGTGGATCAAATGTTTAAAT AGGAAGAACTTGGCAAGGATCTCTTTGATTGTACTTTGTATGTTCTATTGAAATATGATG ATTTTAATGCTGACAAGCACCTGGCTCTTGAAGAATTTTATAGAGCATTCCAAGTGATCC AGTTGAGTCTGCCAGAAGATCAGAAACTAAGCATCACTGCAGCAACTGTGGGACAAAGTG CTGTTCTGAGCTGTGCCATTCAAGGAACCCTGAGACCTCCCATTATCTGGAAAAGGAACA ATATTATTCTAAATAATTTAGATTTGGAAGACATCAATGACTTTGGAGATGATGGGTCCT TGTATATTACTAAGGTTACCACAACTCACGTTGGCAATTACACCTGCTATGCAGATGGCT ATGAACAAGTCTATCAGACTCACATCTTCCAAGTGAATGTTCCTCCAGTCATCC

FIGURE 12

MFKCWSVVLVLGFIFLESEGRPTKEGGYGLKSYQPLMRLRHKQEKNQESSRVKGFMIQDE PFGSCENKYCGLGRHCVTSRETGQAECACMDLCKRHYKPVCGSDGEFYENHCEVHRAAGL KKQKITIVHNEDCFFKGDKCKTTEYSKMKNMLLDLQNQKYIMQENENPNGDDISRKKLLV DQMFKYFDADSNGLVDINELTQVIKQEELGKDLFDCTLYVLLKYDDFNADKHLALEEFYR AFQVIQLSLPEDGKLSITAATYGQSAVUSCAIQGTLRPPIIWKRNNIILNNLDLEDINDF GDDGSLYITKVTTTHVGNYTCYADGYEQVYQTHIFQVNVPPVI

Signal sequence: Amino acids 1-20

N-glycosylation site: Amino acids 318-322

Tyrosine kinase phosphorylation sites: Amino acids 21-29;211-220

N-myristoylation sites: Amino acids 63-69;83-89;317-323

Prokaryotic membrane lipoprotein lipid attachment site: Amino acids 260-271

FIGURE 13

TGCCGGGCTGCGGGGCGCCTTGACTCTCCCTCCACCCTGCCTCCTCGGGCTCCACTCGTC GCAGCTTCCCGCGTCTCCGGCGCAGCTTCTCAGCGGACGACCCTCTCGCTCCGGGGCTGA GCCCAGTCCCTGGATGTTGCTGAAACTCTCGAGATCATGCGCGGGTTTGGCTGCTTC TAGCCCGCTGCCCGGCCCCCGGATCCTGTGTTCCTCGGAAGCCGTTTGCTGCTGCAGAG TTGCACGAACTAGTCATGGTGCTGTGGGAGTCCCCGCGGCAGTGCAGCAGCTGGACACTT GTGA AGCTCGCTGCTTTCCCTACCTCCTTA AGTGACTGCCAAACGCCCACCGGCTGGAAT TGCTCTGGTTATGATGACAGAGAAAATGATCTCTTCCTCTGTGACACCAACACCTGTAAA TTTGATGGGGAATGTTTAAGAATTGGAGACACTGTGACTTGCGTCTGTCAGTTCAAGTGC AACAATGACTATGTGCCTGTGTGTGGCTCCAATGGGGAGAGCTACCAGAATGAGTGTTAC CTGCGACAGGCTGCATGCAAACAGCAGAGTGAGATACTTGTGGTGTCAGAAGGATCATGT GCCACAGATGCAGGATCAGGATCTGGAGATGGAGTCCATGAAGGCTCTGGAGAAACTAGT CAAAAGGAGACATCCACCTGTGATATTTGCCAGTTTGGTGCAGAATGTGACGAAGATGCC GAGGATGTCTGGTGTGTGTAATATTGACTGTTCTCAAACCAACTTCAATCCCCTCTGC GCTTCTGATGGGAAATCTTATGATAATGCATGCCAAATCAAAGAAGCATCGTGTCAGAAA CAGGAGAAAATTGAAGTCATGTCTTTGGGTCGATGTCAAGATAACACAACTACAACTACT AAGTCTGAAGATGGGCATTATGCAAGAACAGATTATGCAGAGAATGCTAACAAATTAGAA GGGAAGTGTGAGCATTCTATCAATATGCAGGAGCCATCTTGCAGGTGTGATGCTGGTTAT ACTGGACAACACTGTGAAAAAAAGGACTACAGTGTTCTATACGTTGTTCCCGGTCCTGTA CGATTTCAGTATGTCTTAATCGCAGCTGTGATTGGAACAATTCAGATTGCTGTCATCTGT CAAAATACAGGGCACTACAGTTCAGACAATACAACAAGAGCGTCCACGAGGTTAATCTAA AGGGAGCATGTTTCACAGTGGCTGGACTACCGAGAGCTTGGACTACACAATACAGTATTA TAGACAAAAGAATAAGACAAGAGATCTACACATGTTGCCTTGCATTTGTGGTAATCTACA ACATTGTCTTGATGTTTTTTCTGTAATGTAAATAAACTATTTATATCACACAATATAGTT TTTTCTTTCCCATGTATTTGTTATATATATAATAATACTCAGTGATGAG

FIGURE 14

MVLWESPRQCSSWTLCEGFCWLLLLPVMLLIVARPVKLAAFPTSLSDCQTPTGWNCSGY DDRENDLFLCDTNTCKFDGECLRIGDTVTCVCQFKCNNDYVPVCGSNGESYQNECYLRQ AACKQQSEILVVSEGSCATDAGSGSGDGVHEGSGETSQKETSTCDLCQFGAECDEDAED WCVCVNIDCSGTNFNPLCASDGKSYDNACQIKEASCQKQEKIEVMSLGRCQDNTTTTTK SEDGHYARTDYAENANKLEESAREHHIPCPEHYNGFCMHGKCEHSINMQEPSCRCDAGY TGQHCEKKDYSVLYVVPGPVRFQYVLIAAVIGTIQIAVICVVVLCITRKCPRSNRIHRQ KONTGHYSSDNTTRASTRLI

FIGURE 15

GGAGCTCAGCCCAGTGGGCAGTCTGAAGATGGCCAATTACACGCTGGCACCAGAGGATGA ATATGATGTCCTCATAGAAGGTGAACTGGAGAGCGATGAGGCAGAGCAATGTGACAAGTA TGACGCCCAGGCACTCTCAGCCCAGCTGGTGCCATCACTCTGCTCTGCTGTTTTTGTGAT CGGTGTCCTGGACAATCTCCTGGTTGTGCTTATCCTGGTAAAATATAAAGGACTCAAACG CGTGGAAAATATCTATCTTCTAAACTTGGCAGTTTCTAACTTGTGTTTCTTGCTTACCCT GCCCTTCTGGGCTCATGCTGGGGGCGATCCCATGTGTAAAATTCTCATTGGACTGTACTT CGTGGGCCTGTACAGTGAGACATTTTTCAATTGCCTTCTGACTGTGCAAAGGTACCTAGT GTTTTTGCACAAGGGCAACTTTTTCTCAGCCAGGAGGAGGGTGCCCTGTGGCATCATTAC AAGTGTCCTGGCATGGGTAACAGCCATTCTGGCCACTTTGCCTGAATACGTGGTTTATAA ACCTCAGATGGAAGACCAGAAATACAAGTGTGCATTTAGCAGAACTCCCTTCCTGCCAGC TGATGAGACATTCTGGAAGCATTTTCTGACTTTAAAAATGAACATTTCGGTTCTTGTCCT CCCCCTATTTATTTTTACATTTCTCTATGTGCAAATGAGAAAAACACTAAGGTTCAGGGA GGCGCCCTACAATATTGCATTTTTCCTGTCCACTTTCAAAGAACACTTCTCCCTGAGTGA CTGCAAGAGCAGCTACAATCTGGACAAAAGTGTTCACATCACTAAACTCATCGCCACCAC CCACTGCTGCATCAACCCTCTCCTGTATGCGTTTCTTGATGGGACATTTAGCAAATACCT CTGCCGCTGTTTCCATCTGCGTAGTAACACCCCACTTCAACCCAGGGGGCAGTCTGCACA AGGCACATCGAGGGAAGAACCTGACCATTCCACCGAAGTGTAAACTAGCATCCACCAAAT GCAAGAAGAATAAACATGGATTTTCATCTTTCTGCATTATTTCATGTAAATTTTCTACAC ATTTGTATACAAAATCGGATACAGGAAGAAAAGGGAGAGGTGAGCTAACATTTGCTAAGC ACTGAATTTGTCTCAGGCACCGTGCAAGGCTCTTTACAAACGTGAGCTCCTTCGCCTCCT ACCACTTGTCCATAGTGTGGATAGGACTAGTCTCATTTCTCTGAGAAGAAAACTAAGGCG CGGAAATTTGTCTAAGATCACATAACTAGGAAGTGGCAGAACTGATTCTCCAGCCCTGGT AGCATTTGCTCAGAGCCTACGCTTGGTCCAGAACATCAAACTCCAAACCCTGGGGACAAA CGACATGAAATAAATGTATTTTAAAACATCTAAAA

FIGURE 16

MANYTLAPEDEYDVLIEGELESDEAEQCDKYDAQALSAQLVPSLCSAVFVIGVLDNLLVV LILVKYKGLKRVENIYLLNLAVSNLCFLLTLPFWAHAGGDPMCKILIGLYFVGLYSETFF NCLLTVQRYLVFLHKGNFFSARRRVPCGIITSVLAWVTAILATLPEYVVYKPQMEDQKYK CAFSRTPFLPADETFWKHFLTLKMNISVLVLPLFIFTFLYVQMRKTLRFREQRYSLFKLV FAIMVVFLLMWAPYNIAFLSTFKEHFSLSDCKSSYNLDKSVHITKLIATTHCCINPLLY AFLDGTFSKYLCRCFHLRSNTPLQPRGQSAQGTSREEPDHSTEV

Signal sequence:

Transmembrane domain:

41-61, 76-96, 109-129, 147-167, 199-219, 237-257, 285-305

7 transmembrane receptor (rhodopsin family): 55-300

N-glycosylation site: 3-6, 205-208

Tyrosine kinase phosphorylation site: 70-76, 171-179, 228-234

N-myristoylation site: 52-57, 136-141, 148-153

G-protein coupled receptors: 55-85, 96-136, 209-220, 235-254, 292-308

FIGURE 17

CGGACGCGTGGGCGGACGCGTGGGCGGCCCACGGCGCCCCGCGGGCTGGGGCGGTCGCTTC TTCCTTCTCCGTGGCCTACGAGGGTCCCCAGCCTGGGTAAAGATGGCCCCATGGCCCCCG AAGGGCCTAGTCCCAGCTGTGCTCTGGGGCCTCAGCCTCTTCCTCAACCTCCCAGGACCT ATCTGGCTCCAGCCCTCTCCACCTCCCCAGTCTTCTCCCCCGCCTCAGCCCCATCCGTGT CATACCTGCCGGGGACTGGTTGACAGCTTTAACAAGGGCCTGGAGAGAACCATCCGGGAC AACTTTGGAGGTGGAAACACTGCCTGGGAGGAAGAGAATTTGTCCAAATACAAAGACAGT GAGACCCGCCTGGTAGAGGTGCTGGAGGGTGTGTGCAGCAAGTCAGACTTCGAGTGCCAC CGCCTGCTGGAGCTGAGTGAGGAGCTGGTGGAGAGCTGGTGGTTTCACAAGCAGCAGGAG GCCCGGACCTCTTCCAGTGGCTGTGCTCAGATTCCCTGAAGCTCTGCTGCCCCGCAGGC ACCTTCGGGCCCTCCTGCCTTCCCTGTCCTGGGGGAACAGAGAGGCCCTGCGGTGGCTAC GGGCAGTGTGAAGGAGAAGGGACACGAGGGGGGCAGCGGGCACTGTGACTGCCAAGCCGGC TACGGGGGTGAGGCCTGTGGCCAGTGTGGCCTTGGCTACTTTGAGGCAGAACGCAACGCC AGCCATCTGGTATGTTCGGCTTGTTTTGGCCCCTGTGCCCGATGCTCAGGACCTGAGGAA TCAAACTGTTTGCAATGCAAGAAGGGCTGGGCCCTGCATCACCTCAAGTGTGTAGACATT GATGAGTGTGGCACAGAGGGAGCCAACTGTGGAGCTGACCAATTCTGCGTGAACACTGAG GGCTCCTATGAGTGCCGAGACTGTGCCAAGGCCTGCCTAGGCTGCATGGGGGCAGGGCCA GGTCGCTGTAAGAAGTGTAGCCCTGGCTATCAGCAGGTGGGCTCCAAGTGTCTCGATGTG GATGAGTGTGAGACAGAGGTGTGTCCGGGAGAGAACAAGCAGTGTGAAAACACCGAGGGC GGTTATCGCTGCATCTGTGCCGAGGGCTACAAGCAGATGGAAGGCATCTGTGTGAAGGAG CAGATCCCAGAGTCAGCAGGCTTCTTCTCAGAGATGACAGAAGACGAGTTGGTGGTGCTG TTGGTGTTCACCGCCATCTTCATTGGGGCTGTGGCGGCCATGACTGGCTACTGGTTGTCA GAGCGCAGTGACCGTGTGCTGGAGGGCTTCATCAAGGGCAGATAATCGCGGCCACCACCT GTAGGACCTCCTCCCACCCACGCTGCCCCCAGAGCTTGGGCTGCCCTCCTGCTGGACACT GCCCAGGTACCCAGGCCCGGGCAGACAAGGCCCCTGGGGTAAAAAGTAGCCCTGAAGGTG GATACCATGAGCTCTTCACCTGGCGGGGACTGGCAGGCTTCACAATGTGTGAATTTCAAA AGTTTTTCCTTAATGGTGGCTGCTAGAGCTTTGGCCCCTGCTTAGGATTAGGTGGTCCTC ACAGGGTGGGGCCATCACAGCTCCCTCCTGCCAGCTGCATGCTGCCAGTTCCTGTTCTG

FIGURE 18

MAPWPPKGLVPAVLWGLSLFLNLPGPIWLQPSPPPQSSPPPQHPCHTCRGLVDSFNKGL ERTIRDNFGGGNTAWEEENLSKYKDSETRLVEVLEGVCSKSDFECHRLIELSEELVESW HKQQEAPDLFQWLCSDSLKLCCPAGTFGPSCLPCPGGTERPCGGYGQCEGESTRGGSGH CDCQAGYGGEACGCQCLGYFEAERNASHLVCSACFGPCARCSGPEESNCLQCKKGWALHH LKCVDIDEGGTEGANCGADQFCVNTEGSYECRDCAKACLGCMGAGPGRCKKCSPGYQQVG SKCLDVDECETEVCPGENKQCENTEGGYRCICAEGYKQMEGICVKEQIPESAGFFSEMTE DELVVLQQMFFGIIICALATLAAKGDLVFTAIFIGAVAAMTGYWLSERSDRVLEGFIKGR

FIGURE 19

GCCCGGGACTGGCGCAAGGTGCCCAAGCAAGGAAAGAAATAATGAAGAGACACATGTGTT AGCTGCAGCCTTTTGAAACACGCAAGAAGGAAATCAATAGTGTGGACAGGGCTGGAACCT ${\tt TTACCACGCTTGTTGGAGTAGATGAGGAATGGGCTCGTGATTATGCTGACATTCCAGC \textbf{AT} }$ GAATCTGGTAGACCTGTGGTTAACCCGTTCCCTCTCCATGTGTCTCCTCCTACAAAGTTT TGTTCTTATGATACTGTGCTTTCATTCTGCCAGTATGTGTCCCAAGGGCTGTCTTTGTTC TTCCTCTGGGGGTTTAAATGTCACCTGTAGCAATGCAAATCTCAAGGAAATACCTAGAGA TCTTCCTCCTGAAACAGTCTTACTGTATCTGGACTCCAATCAGATCACATCTATTCCCAA TGAAATTTTTAAGGACCTCCATCAACTGAGAGTTCTCAACCTGTCCAAAAATGGCATTGA GTTTATCGATGAGCATGCCTTCAAAGGAGTAGCTGAAACCTTGCAGACTCTGGACTTGTC CGACAATCGGATTCAAAGTGTGCACAAAAATGCCTTCAATAACCTGAAGGCCAGGGCCAG AATTGCCAACAACCCCTGGCACTGCGACTGTACTCTACAGCAAGTTCTGAGGAGCATGGC GTCCAATCATGAGACAGCCCACAACGTGATCTGTAAAACGTCCGTGTTGGATGAACATGC TGGCAGACCATTCCTCAATGCTGCCAACGACGCTGACCTTTGTAACCTCCCTAAAAAAAC TACCGATTATGCCATGCTGGTCACCATGTTTGGCTGGTTCACTATGGTGATCTCATATGT GGTATATTATGTGAGGCAAAATCAGGAGGATGCCCGGAGACACCTCGAATACTTGAAATC CCTGCCAAGCAGGCAGAAGAAAGCAGATGAACCTGATGATATTAGCACTGTGGTA**TAG**TG TCCAAACTGACTGTCATTGAGAAAGAAAGAAAGTAGTTTGCGATTGCAGTAGAAATAAGT CACCCCTTAATTGTACCCCCGATGGTATATTTCTGAGTAAGCTACTATCTGAACATTAGT TAGATCCATCTCACTATTTAATAATGAAATTTATTTTTTTAATTTAAAAGCAAATAAAAG

FIGURE 20

MNLVDLWLTRSLSMCLLLQSFVLMILCFHSASMCPKGCLCSSSGGLNVTCSNANLKEIPR DLPPETVLLYLDSNQITSIPNEIFKDLHQLRVLNLSKNGIBFIDEHAFKGVAETLQTLDL SDNRIQSVHKNAFNNLKARARIANNPWHCDCTLQQVLRSMASNHETAHNVICKTSVLDEH AGRPFLNAANDADLCNLPKKTTDYAMLVTMFGWFTMVISYVVYYVRQNQEDARRHLEYLK SLPSRQKKADEPDDISTVV

Signal sequence: amino acids 1-33

Transmembrane domain: amino acids 205-220

N-glycosylation site: amino acids 47-51, 94-98

cAMP- and cGMP-dependent protein kinase phosphorylation site:

Casein kinase II phosphorylation site: amino acids 162-166, 175-179

N-myristoylation site: amino acids 37-43, 45-51, 110-116

FIGURE 21

CGCCACCACTGCGGCCACCGCCAATGAAACGCCTCCCGCTCCTAGTGGTTTTTTCCACTT TGTTGAATTGTTCCTATACTCAAAATTGCACCAAGACACCTTGTCTCCCAAATGCAAAAT GTGAAATACGCAATGGAATTGAAGCCTGCTATTGCAACATGGGATTTTCAGGAAATGGTG TCACAATTTGTGAAGATGATAATGAATGTGGAAATTTAACTCAGTCCTGTGGCGAAAATG CTAATTGCACTAACACAGAAGGAAGTTATTATTGTATGTGTGTACCTGGCTTCAGATCCA GCAGTAACCAAGACAGGTTTATCACTAATGATGGAACCGTCTGTATAGAAAATGTGAATG CAAACTGCCATTTAGATAATGTCTGTATAGCTGCAAATATTAATAAAACTTTAACAAAAA TCAGATCCATAAAAGAACCTGTGGCTTTGCTACAAGAAGTCTATAGAAATTCTGTGACAG ATCTTTCACCAACAGATATAATTACATATATAGAAATATTAGCTGAATCATCTTCATTAC TAGGTTACAAGAACAACACTATCTCAGCCAAGGACACCCTTTCTAACTCAACTCTTACTG AATTTGTAAAAACCGTGAATAATTTTGTTCAAAGGGATACATTTGTAGTTTGGGACAAGT TATCTGTGAATCATAGGAGAACACATCTTACAAAACTCATGCACACTGTTGAACAAGCTA CTTTAAGGATATCCCAGAGCTTCCAAAAGACCACAGAGTTTGATACAAATTCAACGGATA TAGCTCTCAAAGTTTTCTTTTTTGATTCATATAACATGAAACATATTCATCCTCATATGA ATATGGATGGAGACTACATAAATATATTTCCAAAGAGAAAAGCTGCATATGATTCAAATG GCAATGTTGCAGTTGCATTTTTATATATATAAGAGTATTGGTCCTTTGCTTTCATCATCTG ACAACTTCTTATTGAAACCTCAAAATTATGATAATTCTGAAGAGGGGGAAAGAGTCATAT TAACATTTACATTAAGTCATCGAAAGGTCACAGATAGGTATAGGAGTCTATGTGCATTTT GGAATTACTCACCTGATACCATGAATGGCAGCTGGTCTTCAGAGGGCTGTGAGCTGACAT ACTCAAATGAGACCCACACCTCATGCCGCTGTAATCACCTGACACATTTTGCAATTTTGA TGTCCTCTGGTCCTTCCATTGGTATTAAAGATTATAATATTCTTACAAGGATCACTCAAC TAGGAATAATTATTTCACTGATTTGTCTTGCCATATGCATTTTTACCTTCTGGTTCTTCA GTGAAATTCAAAGCACCAGGACAACAATTCACAAAAATCTTTGCTGTAGCCTATTTCTTG CTGAACTTGTTTTCTTGTTGGGATCAATACAAATACTAATAAGCTCTTCTGTTCAATCA TTGCCGGACTGCTACACTACTTCTTTTTAGCTGCTTTTTGCATGGATGTGCATTGAAGGCA TACATCTCTATCTCATTGTTGTGGGTGTCATCTACAACAAGGGATTTTTGCACAAGAATT TTTATATCTTTGGCTATCTAAGCCCAGCCGTGGTAGTTGGATTTTCGGCAGCACTAGGAT ACAGATATTATGGCACAACCAAAGTATGTTGGCTTAGCACCGAAAACAACTTTATTTGGA GTTTTATAGGACCAGCATGCCTAATCATTCTTGTTAATCTCTTGGCTTTTGGAGTCATCA TAAGGTCTTGTGCAAGAGGAGCCCTCGCTCTTCTGTTCCTTCTCGGCACCACCTGGATCT TTGGGGTTCTCCATGTTGTGCACGCATCAGTGGTTACAGCTTACCTCTTCACAGTCAGCA ATGCTTTCCAGGGGATGTTCATTTTTTTATTCCTGTGTGTTTTTATCTAGAAAGATTCAAG AAGAATATTACAGATTGTTCAAAAATGTCCCCTGTTGTTTTGGATGTTTAAGGTAAACAT AGAGAATGGTGGATAATTACAACTGCACAAAAATAAAAATTCCAAGCTGTGGATGACCAA TGTATAAAAATGACTCATCAAATTATCCAATTATTAACTACTAGACAAAAAGTATTTTAA ATCAGTTTTTCTGTTTATGCTATAGGAACTGTAGATAATAAGGTAAAATTATGTATCATA TAGATATACTATGTTTTTCTATGTGAAATAGTTCTGTCAAAAATAGTATTGCAGATATTT ACACGAGAAGTATATGAATGTCCTGAAGGAAACCACTGGCTTGATATTTCTGTGACTCGT GTTGCCTTTGAAACTAGTCCCCTACCACCTCGGTAATGAGCTCCATTACAGAAAGTGGAA CATAAGAGAATGAAGGGGCAGAATATCAAACAGTGAAAAGGGAATGATAAGATGTATTTT GAATGAACTGTTTTTCTGTAGACTAGCTGAGAAATTGTTGACATAAAATAAAGAATTGA AGAAACACATTTTACCATTTTGTGAATTGTTCTGAACTTAAATGTCCACTAAAACAACTT

FIGURE 22

MKRLPLLVVFSTLLNCSYTQNCTKTPCLPNAKCEIRNGIEACYCNMGFSGNGVTICEDDN
BCCNLTQSCGENANCTNTEGSYYCMCVPGFRSSSNQDRFITNDGTVCIENVNANCHLDNV
CIAANINKTLKLRSIKEPVALLQEVYRNSVTDLSPTDIITTIEILAESSSLLGYKNNTI
SAKDTLSNSTLTEFVKTVNNFVQRDTFVVWDKLSVNHRRTHLTKLMHTVEQATLRISQSF
QKTTEFDTNSTDIALKVPFFDSYNMKHHPHNNMDGDYINIFPKRKAAYDSNGNVAVAFL
YYKSIGPLLSSSNDFLKPQNYDNSEEEERVISVISVSWSSNPPTTYELEKITFTLSHR
KVTDRYRSLCAFWNYSPDTMNGSWSSEGCELTYSNETHTSCRCNHLTHFAILMSSGPSIG
IKDYNILTRITQLGIIISLICLAICIFTFWFFSEIQSTRTTTHKNLCCSLFLAELVFLVG
INTNTNKLPCSIIAGLHYFFLAAFAMMCIEGIHLYLIVVGVIYNKGFLHKNFYIFGYLS
PAVVVGFSAALGYRYYGTTKVCWLSTENNFIWSFIGPACLIILVNLLAFGVIIYKVFRHT
AGLKPEVSCFENIRSCARGALALLFLLGTTWIFGVLHVVHASVVTAYLFTVSNAFQGMFI
FLFLCVLSKKIOEEYYRLFKNVPCCFGCLR

FIGURE 23

CTCCTCTTAACATACTTGCAGCTAAAACTAAATATTGCTGCTTGGGGACCTCCTTCTAGC CTTAAATTTCAGCTCATCACCTTCACCTGCCTTGGTCATGGCTCTGCTATTCTCCTTGAT $\overline{\text{CCTTGCCATTTGCACCAGACCTGGATTCCTAGCGTCTCCATCTGGAGTGCGGCTGGTGGG}$ GGGCCTCCACCGCTGTGAAGGGCGGGTGGAGGTGGAACAGAAAGGCCAGTGGGGCACCGT GTGTGATGACGCCTGGGACATTAAGGACGTGGCTGTGTTGTGCCGGGAGCTGGGCTGTGG AGCTGCCAGCGGAACCCCTAGTGGTATTTTGTATGAGCCACCAGCAGAAAAAGAGCAAAA GGTCCTCATCCAATCAGTCAGTTGCACAGGAACAGAAGATACATTGGCTCAGTGTGAGCA AGAAGAAGTTTATGATTGTTCACATGATGAAGATGCTGGGGCATCGTGTGAGAACCCAGA GGGACGCGTGGAAGTGAAGCACCAGAACCAGTGGTATACCGTGTGCCAGACAGGCTGGAG CCTCCGGGCCGCAAAGGTGGTGTGCCGGCAGCTGGGATGTGGGAGGGCTGTACTGACTCA AAAACGCTGCAACAAGCATGCCTATGGCCGAAAACCCATCTGGCTGAGCCAGATGTCATG CTCAGGACGAGAAGCAACCCTTCAGGATTGCCCTTCTGGGCCTTGGGGGAAGAACACCTG CAACCATGATGAAGACACGTGGGTCGAATGTGAAGATCCCTTTGACTTGAGACTAGTAGG AGGAGACAACCTCTGCTCTGGGCGACTGGAGGTGCTGCACAAGGGCGTATGGGGCTCTGT GAAGTCCCTCTCCCCTCCTTCAGAGACCGGAAATGCTATGGCCCTGGGGTTGGCCGCAT CTGGCTGGATAATGTTCGTTGCTCAGGGGAGGAGCAGTCCCTGGAGCAGTGCCAGCACAG ATTTTGGGGGTTTCACGACTGCACCCACCAGGAAGATGTGGCTGTCATCTGCTCAGTGTA GGTGGGCATCATCTAATCTGTTGAGTGCCTGAATAGAAGAAAAACACAGAAGAAGAGGGAGC ATTTACTGTCTACATGACTGCATGGGATGAACACTGATCTTCTTCTGCCCTTGGACTGGG ACTTATACTTGGTGCCCCTGATTCTCAGGCCTTCAGAGTTGGATCAGAACTTACAACATC AGGTCTAGTTCTCAGGCCATCAGACATAGTTTGGAACTACATCACCACCTTTCCTATGTC TCCACATTGCACACAGCAGATTCCCAGCCTCCATAATTGTGTGTATCAACTACTTAAATA TGTTTCTCTGAAGAACTCTGACAAAATACAGATTTTGGTACTGAAAGAGATTCTAGAGGA ACGGAATTTTAAGGATAAATTTTCTGAATTGGTTATGGGGTTTCTGAAATTGGCTCTATA ATCTAATTAGATATAAAATTCTGGTAACTTTATTTACAATAATAAAGATAGCACTATGTG TTCAAA

FIGURE 24

MALLFSLILAICTRPGFLASPSGVRLVGGLHRCEGRVEVEQKGQWGTVCDDGWDIKDVAV LCRELGCGAASGTPSGILYEPPAEKEQKVLIQSVSCTGTEDTLAQCEQEEVYDCSHDEDG GASCENPESSFSPVPEGVRLADGFGHCKGRVEVKHQNGWYTVCQTGWSLRAAKVVCRQLG CGRAVLTQKRCNKHAYGRKPIWLSQMSCSGREATLQDCPSGPWGKNTCNHDEDTWVECED PFDLRLVGGDNLCSGRLEVLHKGWWGSVCDDNWGEKEDQVVCKQLGCGKSLSPSFRDRKC YGPGVGRIMLDNVRCSGEEOSLEQCQHRFWGFHDCTHQEDVAVICSV

Signal sequence: amino acids 1-15

Casein kinase II phosphorylation site: amino acids 47-51, 97-101, 115-119, 209-213, 214-218, 234-238, 267-271, 294-298, 316-320, 336-340

N-myristoylation site: amino acids 29-35, 43-49, 66-72, 68-74, 72-78, 98-104, 137-143, 180-186, 263-269, 286-292

Amidation site: amino acids 196-200

Speract receptor repeated domain signature: amino acids 29-67, 249-287

FIGURE 25

CGGACGCGTGGGCGTCCGGCGGTCGCAGAGCCAGGAGGCGGAGGCGCGCGGGCCAGCCTG GGCCCAGCCCACACCTTCACCAGGGCCCAGGAGCCACCATGTGGCGATGTCCACTGGGG CTACTGCTGTTGCTGCCGCTGGCTGGCCACTTGGCTCTGGGTGCCCAGCAGGGTCGTGGG CGCCGGGAGCTAGCACCGGGTCTGCACCTGCGGGGCATCCGGGACGCGGGAGGCCGGTAC TGCCAGGAGCAGGACCTGTGCTGCCGCGGCCGTGCCGACGACTGTGCCCTGCCCTACCTG GGCGCCATCTGTTACTGTGACCTCTTCTGCAACCGCACGGTCTCCGACTGCTGCCCTGAC TTCTGGGACTTCTGCCTCGGCGTGCCACCCCCTTTTCCCCCGATCCAAGGATGTATGCAT GGAGGTCGTATCTATCCAGTCTTGGGAACGTACTGGGACAACTGTAACCGTTGCACCTGC CAGGAGAACAGGCAGTGGCATGGTGGATCCAGACATGATCAAAGCCATCAACCAGGGCAA CTATGGCTGGCAGGCTGGGAACCACAGCGCCTTCTGGGGCATGACCCTGGATGAGGGCAT TACAGTGCTGAACCCAGGGGAGGTGCTTCCCACAGCCTTCGAGGGCCTCTGAGAAGTGGCC CAACCTGATTCATGAGCCTCTTGACCAAGGCAACTGTGCAGGCTCCTGGGCCTTCTCCAC AGCAGCTGTGGCATCCGATCGTGTCTCAATCCATTCTCTGGGACACATGACGCCTGTCCT GTCGCCCCAGAACCTGCTGTCTTGTGACACCCACCAGCAGCAGGGCTGCCGCGGTGGGCG TCTCGATGGTGCCTGGTGGTTCCTGCGTCGCCGAGGGGTGGTGTCTGACCACTGCTACCC CTTCTCGGGCCGTGAACGAGACGAGGCTGGCCCTGCGCCCCCTGTATGATGCACAGCCG AGCCATGGGTCGGGGCAAGCGCCAGGCCACTGCCCACTGCCCCAACAGCTATGTTAATAA CAATGACATCTACCAGGTCACTCCTGTCTACCGCCTCGGCTCCAACGACAAGGAGATCAT GAAGGAGCTGATGGAGAATGGCCCTGTCCAAGCCCTCATGGAGGTGCATGAGGACTTCTT CCTATACAAGGGAGGCATCTACAGCCACACGCCAGTGAGCCTTGGGAGGCCAGAGAGATA CCGCCGGCATGGGACCCACTCAGTCAAGATCACAGGATGGGGAGAGGAGACGCTGCCAGA TGGAAGGACGCTCAAATACTGGACTGCGGCCAACTCCTGGGGCCCAGCCTGGGGCGAGAG GGGCCACTTCCGCATCGTGCGCGGCGTCAATGAGTGCGACATCGAGAGCTTCGTGCTGGG CGTCTGGGGCCGCGTGGGCATGGAGGACATGGGTCATCACTGAGGCTGCGGGCACCACGC GGGGTCCGGCCTGGGATCCAGGCTAAGGGCCGGCGGAAGAGGCCCCAATGGGGCGGTGAC CCCAGCCTCGCCCGACAGAGCCCGGGGCGCGCAGGGCGCCAAATCCCGGCGC GGGTTCCGCTGACGCAGCGCCCCGCCTGGGAGCCGCGGGCAGGCGAGACTGGCGGAGCCC CAGGCCTCTGGCGCCCCCACTCAAGACTACCAAAGCCAGGACACCTCAAGTCTCCAGCCC TTGCCCAGGTTGGAGTGCAGTGGCCCATCAGGGCTCACTGTAACCTCCGACTCCTGGGTT CAAGTGACCCTCCCACCTCAGCCTCTCAAGTAGCTGGGACTACAGGTGCACCACCACACC TGGCTAATTTTTGTATTTTTTGTAAAGAGGGGGGTCTCACTGTGTTGCCCAGGCTGGTTT CGAACTCCTGGGCTCAAGCGGTCCACCTGCCTCCGCCTCCCAAAGTGCTGGGATTGCAGG TCTTTTAAAATAAAACCAAAGTATTGATAAAAAAAAA

FIGURE 26

MWRCPLGLLLLPLAGHLALGAQQGRGRRELAPGLHLRGIRDAGGRYCQEQDLCCRGRAD DCALPYLGAICYCDLFCNRTVSDCCPDFWDFCLGVPPPFPPIQGCMHGGRIYPVLGTYWD NCNRCTCOENROWHGGSRHDQSHQPGQLWLAGWEPQRLLGHDPG

N-glycosylation site: amino acids 78-82, 161-165

Casein kinase II phosphorylation site: amino acids 80-84, 117-121, 126-130, 169-173, 205-209, 296-300, 411-415

N-myristoylation site: amino acids 21-27, 39-45, 44-50, 104-110, 160-164, 224-230, 269-275, 378-384, 442-448

Amidation site: amino acids 26-30, 318-322

Eukaryotic thiol (cysteine) proteases histidine active site: amino acids 398-409

FIGURE 27

CCCACGCGTCCGGCAGGTTTTTCTTCAAGCCAAGAAGGACACGGATTGGCTGAAGGAGAA AGTGCAGAGCTTGCAGACACTGGCTGCCAACAACTCTGCGTTGGCCAAAGCCAACAACGA CACCCTGGAGGATATGAACAGCCAGCTCAACTCATTCACAGGTCAGATGGAGAACATCAC CACTATCTCTCAAGCCAACGAGCAGAACCTGAAAGACCTGCAGGACTTACACAAAGATGC AGAGAATAGAACAGCCATCAAGTTCAACCAACTGGAGGAACGCTTCCAGCTCTTTGAGAC GGATATTGTGAACATCATTAGCAATATCAGTTACACAGCCCACCACCTGCGGACGCTGAC CAGCAATCTAAATGAAGTCAGGACCACTTGCACAGATACCTTACCAAACACACAGATGAT CTGACCTCCTTGAATAATACCCTGGCCAACATCCGTTTGGATTCTGTTTCTCTCAGGATG CAACAAGATTTGATGAGGTCGAGGTTAGACACTGAAGTAGCCAACTTATCAGTGATTATG GAAGAAATGAAGCTAGTAGACTCCAAGCATGGTCAGCTCATCAAGAATTTTACAATACTA CAAGGTCCACCGGGCCCCAGGGGTCCAAGAGGTGACAGAGGATCCCAGGGACCCCCTGGC CCAACTGGCAACAAGGGACAGAAAGGAGAGAGAGGGGGAGCCTGGACCACCTGGCCCTGCG GGTGAGAGAGGCCCAATTGGACCAGCTGGTCCCCCCGGAGAGCGTGGCGGCAAAGGATCT AAAGGCTCCCAGGGCCCCAAAGGCTCCCGTGGTTCCCCTGGGAAGCCCGGCCCTCAGGGC CCCAGTGGGGACCCAGGCCCCCGGGCCCACCAGGCAAAGAGGGACTCCCCGGCCCTCAG GGCCCTCCTGGCTTCCAGGGACTTCAGGGCACCGTTGGGGAGCCTGGGGTGCCTGGACCT CGGGGACTGCCAGGCTTGCCTGGGGTACCAGGCATGCCAGGCCCCAAGGGCCCCCCCGGC CCTCCTGGCCCATCAGGAGCGGTGGTGCCCCTGGCCCTGCAGAATGAGCCAACCCCGGCA CCGGAGGACAATAGCTGCCCGCCTCACTGGAAGAACTTCACAGACAAATGCTACTATTTT TCAGTTGAGAAAGAAATTTTTGAGGATGCAAAGCTTTTCTGTGAAGACAAGTCTTCACAT CTTGTTTTCATAAACACTAGAGAGGAACAGCAATGGATAAAAAAACAGATGGTAGGGAGA GGGACATCTCCAGACTACAAAAATTGGAAAGCTGGACAGCCGGATAACTGGGGTCATGGC CATGGGCCAGGAGAAGACTGTGCTGGGTTGATTTATGCTGGGCAGTGGAACGATTTCCAA TGTGAAGACGTCAATAACTTCATTTGCGAAAAAGACAGGGAGACAGTACTGTCATCTGCA TTATAACGGACTGTGATGGGATCACATGAGCAAATTTTCAGCTCTCAAAGGCAAAGGACA TTACTGAAAAAAATTGACAGCTAGTGTTTTTTACCATCCGTCATTACCCAAAGACTTGG GAACTAAAATGTTCCCCAGGGTGATATGCTGATTTTCATTGTGCACATGGACTGAATCAC ATAGATTCTCCTCCGTCAGTAACCGTGCGATTATACAAATTATGTCTTCCAAAGTATGGA ACACTCCAATCAGAAAAAGGTTATCATTGGTCGTTGAGTTATGGGAAGAACTTAAGCATA TACTGTGTAAACAGTGCCATACATTTCTAAAATCCCAAGTGTAGGAAAAATATGCAGACA TACAGATATATAGGCCAACTATTAGTAATAATATGAAATATACTTAAAGAGCTTTTAAAA CTTTGTATTTTTGTACAAAAAAAAA

FIGURE 28

MQQDLMRSRLDTEVANLSVIMEEMKLVDSKHGQLIKMFTILQGPPGPRGPRGDRGSQGPP GPTGNKGQKGEKGEPGPPGPAGERGPIGPAGPPGERGGKGSKGSQGPKGSRGS PGKPGPP GPSGDPGPPGPGKEGLPGPQGPGFGGLQGTVGEPGVPGPRGLPGLPGVPGMPGPKGPP GPPGPSGAVVPLALQNEPTPAPEDNSCPPHWKNFTDKCYYFSVEKEIFEDAKLFCEDKSS HLVFINTREEQQWIKKQNVGRESHWIGLTDSERENEWKWLDGTSPDYKNWKAGQPDNWGH GHGPGEDCAGLIYAGOWNDFOCEDVNNFICEKDRETVLSSAL

Signal sequence:

Transmembrane domain:

N-glycosylation site: 16-19, 37-40, 213-216

Tyrosine kinase phosphorylation site: 212-220

N-myristoylation site: 97-102, 100-105, 148-153, 267-272, 293-298, 310-315

Cell attachment sequence: 51-53

C-type lectin domain signature: 308-330

Lectin C-type domain: 233-330

Collagen triple helix repeat: 43-102, 127-186

FIGURE 29

GGACTAATCTGTGGGAGCAGTTTATTCCAGTATCACCCAGGGTGCAGCCACACCAGGACT AGTGTCTGAGAACATTTACATTATAGATAAGTAGTACATGGTGGATAACTTCTACTTTTA GGAGGACTACTCTCTGCACAGTCCTAGACTGGTCTTCTACACTAAGACACCATGAAGG AGTATGTGCTCCTATTATTCCTGGCTTTGTGCTCTGCCAAACCCTTCTTAGCCCTTCAC ACATCGCACTGAAGAATATGATGCTGAAGGATATGGAAGACACAGATGATGATGATGATG ATGATGATGATGATGATGATGAGGGACAACTCTCTTTTTCCAACAAGAGAGCCAAGAA GCCATTTTTTTCCATTTGATCTGTTTCCAATGTGTCCATTTGGATGTCAGTGCTATTCAC CTCGAATGCTTGATCTTCAAAACAATAAAATTAAGGAAATCAAAGAAAATGATTTTAAAG GACTCACTTCACTTTATGGTCTGATCCTGAACAACAACAAGCTAACGAAGATTCACCCAA AAGCCTTTCTAACCACAAAGAAGTTGCGAAGGCTGTATCTGTCCCACAATCAACTAAGTG AAATACCACTTAATCTTCCCAAATCATTAGCAGAACTCAGAATTCATGAAAATAAAGTTA AGAAAATACAAAAGGACACATTCAAAGGAATGAATGCTTTACACGTTTTGGAAATGAGTG CAAACCCTCTTGATAATAATGGGATAGAGCCAGGGGCATTTGAAGGGGTGACGGTGTTCC ATATCAGAATTGCAGAAGCAAAACTGACCTCAGTTCCTAAAGGCTTACCACCAACTTTAT TGGAGCTTCACTTAGATTATAATAAAATTTCAACAGTGGAACTTGAGGATTTTAAACGAT ACAAAGAACTACAAAGGCTGGGCCTAGGAAACAACAAAATCACAGATATCGAAAATGGGA TTGCAAGAGTGGGAGTAAATGACTTCTGTCCAACAGTGCCAAAGATGAAGAAATCTTTAT ACAGTGCAATAAGTTTATTCAACAACCCGGTGAAATACTGGGAAATGCAACCTGCAACAT ${\tt TTCGTTGTGTTTTGAGCAGAATGAGTGTTCAGCTTGGGAACTTTGGAATGTAATAATTAG}$ TAATTGGTAATGTCCATTTAATATAAGATTCAAAAATCCCTACATTTGGAATACTTGAAC TCTATTAATAATGGTAGTATTATATATACAAGCAAATATCTATTCTCAAGTGGTAAGTCC ACTGACTTATTTTATGACAAGAAATTTCAACGGAATTTTGCCAAACTATTGATACATAAG GGGTTGAGAGAAACAAGCATCTATTGCAGTTTCCTTTTTGCGTACAAATGATCTTACATA AATCTCATGCTTGACCATTCCTTTCTTCATAACAAAAAAGTAAGATATTCGGTATTTAAC AAAATTTGTGCTCTTTCATTTGCTGTTAGAAAAACAGAATTAACAAAGACAGTAATGTGA AGAGTGCATTACACTATTCTTATTCTTTAGTAACTTGGGTAGTACTGTAATATTTTTAAT CATCTTAAAGTATGATTTGATATAATCTTATTGAAATTACCTTATCATGTCTTAGAGCCC GTCTTTATGTTTAAAACTAATTTCTTAAAATAAAGCCTTCAGTAAATGTTCATTACCAAC ATTATTACCTGATTTAAAAATCTCTGTAAAAACGTGTAGTGTTTCATAAAATCTGTAACT CGCATTTTAATGATCCGCTATTATAAGCTTTTAATAGCATGAAAATTGTTAGGCTATATA GAAGAGCCTGGACACTAACAATTCTACACCAAATTGTCTCTTCAAATACGTATGGACTGG ATAACTCTGAGAAACACATCTAGTATAACTGAATAAGCAGAGCATCAAATTAAACAGACA GAAACCGAAAGCTCTATATAAATGCTCAGAGTTCTTTATGTATTTCTTATTGGCATTCAA CATATGTAAAATCAGAAAACAGGGAAATTTTCATTAAAAATATTGGTTTGAAAT

FIGURE 30

MKEYVLLLFLALCSAKPFFSPSHIALKNMMLKDMEDTDDDDDDDDDDDDDDDDDDDDLNSLFPTRE PRSHFFPPDLFPMCPFGCQCYSRVVHCSDLGLTSVPTNIPFDTRMLDLQNNKIKEIKEND FKGLTSLYGLILNNNKLTKIHPKAFLTTKKLRRLYLSHNQLSEIPLNLPKSLAELRIHEN KVKKIQKDTFKGMNALHVLEMSANPLDNNGIEPGAFEGVTVFHIRIAEAKLTSVPKGLPP TLLELHLDYNKISTVELEDFKRYKELQRLGLGNNKITDIENGSLANIPRVREIHLENNKL KKIPSGLPELKYLQIIFLHSNSIARVGVNDFCPTVPKMKKSLYSAISLFNNPVKYWEMQP ATFRCVLSRMSVOLGNFGM

Signal sequence: amino acids 1-15

N-glycosylation site: amino acids 281-285

N-myristoylation sites: amino acids 129-135, 210-216, 214-220, 237-243, 270-276, 282-288

Leucine zipper pattern: amino acids 154-176

FIGURE 31

GTTCCAGAACTCTCCATCCGGACTAGTTATTGAGCATCTGCCTCTCATATCACCAGTGGC CATCTGAGGTGTTTCCCTGGCTCTGAAGGGGTAGGCACG**ATG**GCCAGGTGCTTCAGCCTG GTGTTGCTTCTCACTTCCATCTGGACCACGAGGCTCCTGGTCCAAGGCTCTTTGCGTGCA GAAGAGCTTTCCATCCAGGTGTCATGCAGAATTATGGGGATCACCCTTGTGAGCAAAAAG GCGAACCAGCAGCTGAATTTCACAGAAGCTAAGGAGGCCTGTAGGCTGCTGGGACTAAGT GGCTGGGTTGGAGATGGATTCGTGGTCATCTCTAGGATTAGCCCAAACCCCAAGTGTGGG AAAAATGGGGTGGGTGTCCTGATTTGGAAGGTTCCAGTGAGCCGACAGTTTGCAGCCTAT TGTTACAACTCATCTGATACTTGGACTAACTCGTGCATTCCAGAAATTATCACCACCAAA GATCCCATATTCAACACTCAAACTGCAACACAAACAACAGAATTTATTGTCAGTGACAGT ACCTACTCGGTGGCATCCCCTTACTCTACAATACCTGCCCCTACTACTACTCCTCCTGCT CCAGCTTCCACTTCTATTCCACGGAGAAAAAAATTGATTTGTCTCACAGAAGTTTTTATG GAAACTAGCACCATGTCTACAGAAACTGAACCATTTGTTGAAAATAAAGCAGCATTCAAG TTTGGTGCTGCAGCTGGTCTTGGATTTTGCTATGTCAAAAGGTATGTGAAGGCCTTCCCT TTTACAAACAAGAATCAGCAGAAGGAAATGATCGAAACCAAAGTAGTAAAGGAGGAGAAG GCCAATGATAGCAACCCTAATGAGGAATCAAAGAAAACTGATAAAAACCCAGAAGAGTCC AAGAGTCCAAGCAAAACTACCGTGCGATGCCTGGAAGCTGAAGTTTAGATGACAGAAA $\mathsf{TGAGGAGACACCTGAGGCTGGTTTCTTTCATGCTCCTTACCCTGCCCCAGCTGGGGAA}$ ATCAAAAGGGCCAAAGAACCAAAGAAGAAGTCCACCCTTGGTTCCTAACTGGAATCAGC CCTGTCTGGATCCTATCCTCCTACCTCCAAAGCTTCCCACGGCCTTTCTAGCCTGGCTAT GTCCTAATAATATCCCACTGGGAGAAAGGAGTTTTGCAAAGTGCAAGGACCTAAAACATC TCATCAGTATCCAGTGGTAAAAAGGCCTCCTGGCTGTCTGAGGCTAGGTGGGTTGAAAGC CAAGGAGTCACTGAGACCAAGGCTTTCTCTACTGATTCCGCAGCTCAGACCCTTTCTTCA GCTCTGAAAGAGAAACACGTATCCCACCTGACATGTCCTTCTGAGCCCGGTAAGAGCAAA AGAATGCCAGAAAAGTTTAGCCCCTGAAAGCCATGGAGATTCTCATAACTTGAGACCTAA TCTCTGTAAAGCTAAAATAAAGAAATAGAACAAGGCTGAGGATACGACAGTACACTGTCA GCAGGGACTGTAAACACAGACAGGGTCAAAGTGTTTTCTCTGAACACATTGAGTTGGAAT CACTGTTTAGAACACACACTTACTTTTTCTGGTCTCTACCACTGCTGATATTTTCTCT AGGAAATATACTTTTACAAGTAACAAAAATAAAAACTCTTATAAATTTCTATTTTTATCT GAGTTACAGAAATGATTACTAAGGAAGATTACTCAGTAATTTGTTTAAAAAGTAATAAAA TTCAACAAACATTTGCTGAATAGCTACTATATGTCAAGTGCTGTGCAAGGTATTACACTC TGTAATTGAATATTATTCCTCAAAAAATTGCACATAGTAGAACGCTATCTGGGAAGCTAT TTTTTCAGTTTTGATATTTCTAGCTTATCTACTTCCAAACTAATTTTTATTTTTGCTGA GACTAATCTTATTCATTTTCTCTAATATGGCAACCATTATAACCTTAATTTATTAAC ATACCTAAGAAGTACATTGTTACCTCTATATACCAAAGCACATTTTAAAAGTGCCATTAA CAAATGTATCACTAGCCCTCTTTTTCCAACAAGAAGGGACTGAGAGATGCAGAAATATT TGTGACAAAAATTAAAGCATTTAGAAAACTT

FIGURE 32

MARCFSLVLLLTSIWTTRLLVQGSLRAEELSIQVSCRIMGITLVSKKANQQLNFTEAKEA
CRLLGLSLAGKDQVETALKASFETCSYGWVGDGFVVISRISPNPKCGKNGVGVLIWKVPV
SRQFAAYCNSSDTWTNSCIPEIITTKDPIFNTQTATQTTEFIVSDSTYSVASPYSTIPA
PTTTPPAPASTSIPRRKKLICVTEVFMETSTMSTETEPFVENKAAFKMEAAGFGGVPTAL
LVLALLFFGAAAGLGFCYVKRYVKAFPFTNKNQQKEMIETKVVKEEKANDSNPNEESKKT
DKNPEESKSPSKTTYRCLEAEV

Signal sequence:

Transmembrane domain: amino acids 235-254

N-glycosylation site: amino acids 53-57, 130-134, 289-293

Casein kinase II phosphorylation site: amino acids 145-149, 214-218

Tyrosine kinase phosphorylation site: amino acids 79-88

N-myristoylation site: amino acids 23-29, 65-71, 234-240, 235-239, 249-255, 253-259

FIGURE 33

GAAAAAAAAAAAAAGGGAAGCAAGCTTAGCTGTACACCCTGAGTCTTGCAAAAGCTGCAG CCCCACCCAGGAGCAGGGTGGTGGCTGGGGCGATGGTGGACGCCCTGAAGATGTCCC<u>ATG</u> GCTACTGAAGGGGCTGCCCAGTTAGGGAACAGAGTGGCGGGCATGGTGTAGCCTATGG GTGCTGCTCCTGGTGTCTTCAGTTCTGGCTCTGGAAGAGGTATTGCTGGACACCACCGGA GAGACATCTGAGATTGGCTGGCTCACCTACCCACCAGGGGGGTGGGACGAGGTGAGTGTT CTGGACGACCAGCGACGCCTGACTCGGACCTTTGAGGCATGTCATGTGGCAGGGGCCCCT CCAGGCACCGGGCAGGACAATTGGTTGCAGACACACTTTGTGGAGCGCGCGGGGCCCAG AGGGCGCACATTCGACTCCACTTCTCTGTGCGGGCATGCTCCAGCCTGGGTGTGAGCGGC GGCACCTGCCGGGAGACCTTCACCCTTTACTACCGTCAGGCTGAGGAGCCCGACAGCCCT GACAGCGTTTCCTCCTGGCACCTCAAACGTTGGACCAAGGTGGACACAATTGCAGCAGAC GAGAGCTTTCCATCCTCCTCCTCCTCCTCCTCCTCTTCTTCTTCCTCTGCAGCGTGGGCT GTGGGACCCCACGGGGCTGGGCAGCGGCTGGACTGCAACTGAACGTCAAAGAGCGGAGC GCCCTGGTCGCTGTCAGGCTCTTCTCCTACACCTGCCCTGCCGTGCTCCGATCCTTTGCT TCCTTTCCAGAGACGCAGGCCAGTGGGGGCTGGGGGGGCCTCCCTGGTGGCAGCTGTGGGC CCCCCAGGCTGCACTGCAACGGGGAGGGCAAGTGGATGGTAGCTGTCGGGGGCTGCCGC TGCCAGCCTGGATACCAACCAGCACGAGGAGACAAGGCCTGCCCAAGCCTGCCCACGGGGG CTCTATAAGGCTTCTGCTGGGAATGCTCCCTGCTCACCATGCCCTGCCCGCAGTCACGCT CCCAACCCAGCAGCCCCCGTTTGCCCCTGCCTGGAGGGCTTCTACCGGGCCAGTTCCGAC CCACCAGAGGCCCCCTGCACTGGTCCTCCATCGGCTCCCCAGGAGCTTTGGTTTGAGGTG CAAGGCTCAGCACTCATGCTACACTGGCGCCTGCCTCGGGAGCTGGGGGGTCGAGGGGAC CTGCTCTTCAATGTCGTGTGCAAGGAGTGTGAAGGCCGCCAGGAACCTGCCAGCGGTGGT GGGGGCACTTGTCACCGCTGCAGGGATGAGGTCCACTTCGACCCTCGCCAGAGAGGCCTG ACTGAGAGCCGAGTGTTAGTGGGGGGACTCCGGGCACACGTACCCTACATCTTAGAGGTG CAGGCTGTTAATGGGGTGTCTGAGCTCAGCCCTGACCCTCCTCAGGCTGCAGCCATCAAT GTCAGCACCAGCCATGAAGTGCCCTCTGCTGTCCCTGTGGTGCACCAGGTGAGCCGGGCA TCCAACAGCATCACGGTGTCCTGGCCGCAGCCCGACCAGACCAATGGGAACATCCTGGAC TATCAGCTCCGCTACTATGACCAGGCAGAAGACGAATCCCACTCCTTCACCCTGACCAGC GAGACCAACACTGCCACCGTGACACAGCTGAGCCCTGGCCACATCTATGGTTTCCAGGTG CGGGCCGGACTGCTGCCGGCCACGGCCCCTACGGGGGCAAAGTCTATTTCCAGACACTT CCTCAAGGGGAGCTGTCTTCCCAGCTTCCGGAAAGACTCTCCTTGGTGATCGGCTCCACC CTGGGGGCTTTGGCCTTCCTGCTGGCAGCCATCACCGTGCTGGCGGTCGTCTTCCAG CGGAAGCGGCGTGGGACTGGCTACACGGAGCAGCTGCAGCAATACAGCAGCCCAGGACTC GGGGTGAAGTATTACATCGACCCCTCCACCTACGAGGACCCCTGTCAGGCCATCCGAGAA CTTGCCCGGGAAGTCGATCCTGCTTATATCAAGATTGAGGAGGTCATTGGGACAGGCTCT ATCCAGGCCCTGTGGGCCGGGGGCGCCGAAAGCCTGCAGATGACCTTCCTGGGCCGGGCC GCAGTGCTGGGTCAGTTCCAGCACCCCAACATCCTGCGGCTGGAGGGCGTGGTCACCAAG AGCCGACCCCTCATGGTGCTGACGGAGTTCATGGAGCTTGGCCCCCTGGACAGCTTCCTC AGGCAGCGGGAGGGCCAGTTCAGCAGCCTGCAGCTGGTGGCCATGCAGCGGGGAGTGGCT GTGCTGGTGAATAGCCACTTGGTGTGCAAGGTGGCCCGTCTTGGCCACAGTCCTCAGGGC CCAAGTTGTTTGCTTCGCTGGGCAGCCCCAGAGGTCATTGCACATGGAAAGCATACTCAT GTGGGAAGTGATGAGTTATGGAGAACGGCCTTACTGGGACATGAG<u>TGA</u>GCAGGAGGTACT AAATGCAATAGAGCAGGAGTTCCGGCTGCCCCCGCCTCCAGGCTGTCCTCCTGGATTACA

FIGURE 34

MATEGAAQLGNRVAGMVCSLWVLLLVSSVLALEEVLLDTTGETSEIGWLTYPPGGWDEVS
VLDDQRRLTRTFEACHVAGAPPGTGQDNWLQTHFVERRAGRAHIRLHFSVRACSSLGVS
GGTCRETFILYYRQAEEPDSPDSVSWHLKRWTKVDTIADBESFPSSSSSSSSSAW
AVGPHGAGQRAGLQLNVKERSFGPLTQRGFYVAFQDTGACLALVAVRLFSYTCPAVLRSF
ASFPETQASGAGGASLVAAVGTCVAHAEPEEDGVGGQAGGSPPRLHCNGEGKWMVANGG
RCQPGYQPARGDKACQACPRGLYKASAGNAPCSPCPARSHAPNPAAPVCPCLEGFYRASS
DPPEAPCTGPPSAPQELWFEVQGSALMLHWRLPRELGGRGDLLFNVVCKECEGRQEPASG
GGGTCHRCRDEVHFDPRQRGLTESRVLVGGLRAHVPYILEVQANNGVSELSFDPPQAAASI
NVSTSHEVPSAVPVYUQVSRASNSITVSWPQPDOTNGNILDYGLRYYDQAEDESHSFTLT
SETNTATVTQLSPGHIYGFQVRARTAGHGPYGGKVYFQTLPQGELSSQLPERLSLVIGS
TLGALAFLLLAAITVLAVVFQRKRRGTGYTEQLQQYSSPGLGVKYYIDPSTYEDPCQAIS
ELARSVDPAYIKIESVIGTGSFGEVRQGRLQPRGRREDTVATQALWAGGAESLQMTFLGR
AAVLGQFQHPNILRLEGVVTKSRPLMVLTEFMELGPLDSFLRQREGGFSSLQLVAMQRGV
AAAMQYLSSFAFVHRSLSAHSVLVNSHLVCKVARLGHSPQGPSCLLRWAAPEVIAHGKHT

signal sequence: Amino acids 1-31

Transmembrane domains: Amino acids 217-234;598-618

N-glycosylation site: Amino acids 481-485

Glycosaminoglycan attachment sites: Amino acids 249-253;419-423

 $\mathtt{cAMP-}$ and $\mathtt{cGMP-} \mathtt{dependent}$ protein kinase phosphorylation sites:

Amino acids 66-70;150-154;624-628

Tyrosine kinase phosphorylation sites: Amino acids 644-673;664-671

N-myristovlation sites:

Amino acids 10-16;15-21;79-85;99-105;118-124;188-194; 192-198;218-224;250-256;261-267;275-281;276-282;298-304;321-327;328-334;420-426;421-427;440-446;449-455;599-605;626-632; 708-714:766-772;779-785

Amidation site: Amino acids 693-697

Cell attachment sequences:
Amino acids 310-313:399-402

FIGURE 35

GCATCCGCAGGTTCCCGCGGACTTGGGGGCGCCCGCTGAGCCCCGGCGCCCCGCAGAAGAC TTGTGTTTGCCTCCTGCAGCCTCAACCCGGAGGGCAGCGAGGGCCTACCACCATGATCAC TGGTGTGTTCAGCATGCGCTTGTGGACCCCAGTGGGCGTCCTGACCTCGCTGGCGTACTG CCTGCACCAGCGGCGGGTGGCCCTGGCCGAGCTGCAGGAGGCCGATGGCCAGTGTCCGGT CGACCGCAGCCTGCTGAAGTTGAAAATGGTGCAGGTCGTGTTTCGACACGGGGCTCGGAG TCCTCTCAAGCCGCTCCCGCTGGAGGAGCAGGTAGAGTGGAACCCCCAGCTATTAGAGGT CCCACCCCAAACTCAGTTTGATTACACAGTCACCAATCTAGCTGGTGGTCCGAAACCATA TTCTCCTTACGACTCTCAATACCATGAGACCACCCTGAAGGGGGGCATGTTTGCTGGGCA TGTGGAAGACATTCCCTTTCTTTCACCAACCTTCAACCCACAGGAGGTCTTTATTCGTTC TCAGAAAGAAGACCCATCATCATCCACACTGATGAAGCAGATTCAGAAGTCTTGTATCC CAACTACCAAAGCTGCTGGAGCCTGAGGCAGAGAACCAGAGGCCGGAGGCAGACTGCCTC TTTACAGCCAGGAATCTCAGAGGATTTGAAAAAGGTGAAGGACAGGATGGGCATTGACAG TAGTGATAAAGTGGACTTCTTCATCCTCCTGGACAACGTGGCTGCCGAGCAGGCACAAA CCTCCCAAGCTGCCCCATGCTGAAGAGATTTGCACGGATGATCGAACAGAGAGCTGTGGA CACATCCTTGTACATACTGCCCAAGGAAGACAGGGAAAGTCTTCAGATGGCAGTAGGCCC ATTCCTCCACATCCTAGAGAGCAACCTGCTGAAAGCCATGGACTCTGCCACTGCCCCCGA CAAGATCAGAAAGCTGTATCTCTATGCGGCTCATGATGTGACCTTCATACCGCTCTTAAT GACCCTGGGGATTTTTGACCACAAATGGCCACCGTTTGCTGTTGACCTGACCATGGAACT TTACCAGCACCTGGAATCTAAGGAGTGGTTTGTGCAGCTCTATTACCACGGGAAGGAGCA GGTGCCGAGAGGTTGCCCTGATGGGCTCTGCCCGCTGGACATGTTCTTGAATGCCATGTC AGTTTATACCTTAAGCCCAGAAAAATACCATGCACTCTGCTCTCAAACTCAGGTGATGGA AGTTGGAAATGAAGAGTAACTGATTTATAAAAGCAGGATGTGTTGATTTTAAAATAAAGT CCCTTTATACAATG

FIGURE 36

MITGVFSMRLWTPVGVLTSLAYCLHQRRVALAELQEADGQCPVDRSLLKLKMVQVVFRHG ARSPLKPLPLEEQVEWNPQLLEVPPQTOFDYTVTNLAGGPKPYSPYDSQYHETTLKGGMF AGQITKVGMQQMFALGERLRKNYVEDIPFLSPTFNPQEVFIRSTNIFRNLESTRCLLAGL FQCQKEGPIIIHTDEADSEVLYPNYQSCWSLRQRTRGRRQTASLQPGISEDLKKVKDRMG IDSSDKVDFFILLDNVAAEQAHNLPSCPMLKRFARMIEQRAVDTSLYILPKEDRESLQMA VGPFLHILESNLLKAMDSATAPDKIRKLYLYAAHDVTFIPLLMTLGIFDHKWPPFAVDLT MELYQHLESKWFVQLYYHGKEQVPRGCPDGLCPLDMFLNAMSVYTLSPEKYHALCSQTQ VMEVGNEE

Signal sequence: amino acids 1-23

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 218-222

Casein kinase II phosphorylation site: amino acids 87-91, 104-108, 320-324

Tyrosine kinase phosphorylation site: amino acids 280-288

N-myristoylation site: amino acids 15-21, 117-123, 118-124, 179-185, 240-246, 387-393

Amidation site: amino acids 216-220

Leucine zipper pattern: amino acids 10-32

Histidine acid phosphatases phosphohistidine signature: amino acids 50-65

FIGURE 37

ACTGCACTCGGTTCTATCGATTGAATTCCCCGGGGATCCTCTAGAGATCCCTCGACCTCG GAAGGTGAAGGCCATGGACTTCATCACCTCCACAGCCATCCTGCCCCTGCTGTTCGGCTG CCTGGGCGTCTTCGGCCTCTTCCGGCTGCTGCAGTGCGCGCGGGAAGGCCTACCTGCG GAATGCTGTGGTGGTGATCACAGGCGCCACCTCAGGGCTGGGCAAAGAATGTGCAAAAGT CTTCTATGCTGCGGGTGCTAAACTGGTGCTCTGTGGCCGGAATGGTGGGGCCCTAGAAGA GCTCATCAGAGAACTTACCGCTTCTCATGCCACCAAGGTGCAGACACACAAGCCTTACTT GGTGACCTTCGACCTCACAGACTCTGGGGCCATAGTTGCAGCAGCAGCTGAGATCCTGCA GTGCTTTGGCTATGTCGACATACTTGTCAACAATGCTGGGATCAGCTACCGTGGTACCAT CATGGACACCACAGTGGATGTGGACAAGAGGGTCATGGAGACAAACTACTTTGGCCCAGT TGCTCTAACGAAAGCACTCCTGCCCTCCATGATCAAGAGGAGGCAAGGCCACATTGTCGC CATCAGCAGCATCCAGGGCAAGATGAGCATTCCTTTTCGATCAGCATATGCAGCCTCCAA GCACGCAACCCAGGCTTTCTTTGACTGTCTGCGTGCCGAGATGGAACAGTATGAAATTGA GGTGACCGTCATCAGCCCCGGCTACATCCACCACCAACCTCTCTGTAAATGCCATCACCGC GGATGGATCTAGGTATGGAGTTATGGACACCACCACAGCCCAGGGCCGAAGCCCTGTGGA GGTGGCCCAGGATGTTCTTGCTGCTGTGGGGAAGAAGAAGAAGATGTGATCCTGGCTGA CTTACTGCCTTCCTTGGCTGTTTATCTTCGAACTCTGGCTCCTGGGCTCTTCTTCAGCCT CATGGCCTCCAGGGCCAGAAAAGAGCGGAAATCCAAGAACTCCTAGTACTCTGACCAGCC AGGGCCAGGGCAGAGAAGCACTCTTAGGCTTGCTTACTCTACAAGGGACAGTTGCAT TTGTTGAGACTTTAATGGAGATTTGTCTCACAAGTGGGAAAGACTGAAGAAACACATCTC GTGCAGATCTGCTGGCAGAGGACAATCAAAAACGACAACAAGCTTCTTCCCAGGGTGAGG GGAAACACTTAAGGAATAAATATGGAGCTGGGGTTTAACACTAAAAACTAGAAATAAACA TCTCAAACAGTAAAAAAAAAAAAAAGGGCGGCCGCGACTCTAGAGTCGACCTGCAGAAG CTTGGCCGCCATGGCCCAACTTGTTTATTGCAGCTTATAATGGTTAC

FIGURE 38

MDFITSTAILPLLFGCLGVFGLFRLLQWVRGKAYLRNAVVVITGATSGLGKECAKVFYAA
GAKLVLCGRNGGALEELIRELTASHATKVQTHKPYLVTFDLTDSGAIVAAAAEILQCFGY
VDILVNNAGISYRGTIMDTTVDVDVKRVMETNYFGPVALTKALLPSMIKRRQGHIVAISSI
QGKMSIPFRSAYAASKHATQAFFDCLRAEMEQYEIEVTVISPGYIHTNLSVNAITADGSR
YGVMDTTTAQGRSPVEVAQDVLAAVGKKKKDVILADLLPSLAVYLRTLAPGLFFSLMASR
ARKERKSKNS

Signal sequence:

Transmembrane domain: amino acids 104-120, 278-292

N-glycosylation site: amino acids 228-232

Glycosaminoglycan attachment site: amino acids 47-51

Casein kinase II phosphorylation site: amino acids 135-139, 139-143, 253-257

Tyrosine kinase phosphorylation site: amino acids 145-153, 146-153

N-myristoylation site: amino acids 44-50, 105-111, 238-244, 242-248, 291-297

Amidation site: amino acids 265-269

Prokaryotic membrane lipoprotein lipid attachment site: amino acids 6-17

FIGURE 39

GCAAGCCAAGGCGCTGTTTGAGAAGGTGAAGAAGTTCCGGACCCATGTGGAGGAGGGGGACATTGT GTACCGCCTCTACATGCGGCAGACCATCATCAAGGTGATCAAGTTCATCCTCATCATCTGCTACAC CTACCGCACCTACCGCTGTGCCCACCCCCTGGCCACACTCTTCAAGATCCTGGCGTCCTTCTACAT CAGCCTAGTCATCTTCTACGGCCTCATCTGCATGTACACACTGTGGTGGATGCTACGGCGCTCCCT CAAGAAGTACTCGTTTGAGTCGATCCGTGAGGAGGAGCAGCTACAGCGACATCCCCGACGTCAAGAA CGACTTCGCCTTCATGCTGCACCTCATTGACCAATACGACCCGCTCTACTCCAAGCGCTTCGCCGT CTTCCTGTCGGAGGTGAGTGAGAACAAGCTGCGGCAGCTGAACCTCAACAACGAGTGGACGCTGGA CAAGCTCCGGCAGCGGCTCACCAAGAACGCGCAGGACAAGCTGGAGCTGCACCTGTTCATGCTCAG TGGCATCCCTGACACTGTTTTGACCTGGTGGAGCTGGAGGTCCTCAAGCTGGAGCTGATCCCCGA CGTGACCATCCCGCCCAGCATTGCCCAGCTCACGGGCCTCAAGGAGCTGTGGCTCTACCACACAGC GGCCAAGATTGAAGCGCCTGCGCTGGCCTTCCTGCGCGAGAACCTGCGGGCGCTGCACATCAAGTT CACCGACATCAAGGAGATCCCGCTGTGGATCTATAGCCTGAAGACACTGGAGGAGCTGCACCTGAC GGGCAACCTGAGCGCGGAGAACAACCGCTACATCGTCATCGACGGGCTGCGGGAGCTCAAACGCCT CAAGGTGCTGCGGCTCAAGAGCAACCTAAGCAAGCTGCCACAGGTGGTCACAGATGTGGGCGTGCA CCTGCAGAAGCTGTCCATCAACAATGAGGGCACCAAGCTCATCGTCCTCAACAGCCTCAAGAAGAT GGCGAACCTGACTGAGCTGGAGCTGATCCGCTGCGACCTGGAGCGCATCCCCACTCCATCTTCAG CCTCCACAACCTGCAGGAGATTGACCTCAAGGACAACAACCTCAAGACCATCGAGGAGATCATCAG CTTCCAGCACCTGCACCGCCTCACCTGCCTTAAGCTGTGGTACAACCACATCGCCTACATCCCCAT CCAGATCGGCAACCTCACCAACCTGGAGCGCCTCTACCTGAACCGCAACAAGATCGAGAAGATCCC CACCCAGCTCTTCTACTGCCGCAAGCTGCGCTACCTGGACCTCAGCCACAACAACCTGACCTTCCT CCCTGCCGACATCGGCCTCCTGCAGAACCTCCAGAACCTAGCCATCACGGCCAACCGGATCGAGAC GCTCCCTCCGGAGCTCTTCCAGTGCCGGAAGCTGCGGGCCCTGCACCTGGGCAACAACGTGCTGCA GTCACTGCCCTCCAGGGTGGGCGAGCTGACCAACCTGACGCAGATCGAGCTGCGGGGCAACCGGCT GGAGTGCCTGCCTGTGGAGCTGGGCGAGTGCCCACTGCTCAAGCGCAGCGGCTTGGTGGTGGAGGA GGACCTGTTCAACACACTGCCACCCGAGGTGAAGGAGCGGCTGTGGAGGGCTGACAAGGAGCAGGC $\mathtt{C} \underline{\mathbf{TGA}}$ GCGAGGCCGGCCCAGCACAGCAAGCAGCAGGACCGCTGCCCAGTCCTCAGGCCCGGAGGGGC AGGCCTAGCTTCTCCCAGAACTCCCGGACAGCCAGGACAGCCTCGCGGCTGGGCAGGAGCCTGGGG CCGCTTGTGAGTCAGGCCAGAGCGAGAGGACAGTATCTGTGGGGCTGGCCCCTTTTCTCCCTCTGA GACTCACGTCCCCCAGGGCAAGTGCTTGTGGAGGAGGAGCAAGTCTCAAGAGCGCAGTATTTGGATA ATCAGGGTCTCCTCCTGGAGGCCAGCTCTGCCCCAGGGGCTGAGCTGCCACCAGAGGTCCTGGGA CCCTCACTTTAGTTCTTGGTATTTATTTTTCTCCATCTCCCACCTCCTTCATCCAGATAACTTATA CATTCCCAAGAAAGTTCAGCCCAGATGGAAGGTGTTCAGGGAAAGGTGGGCTGCCTTTTCCCCTTG TCCTTATTTAGCGATGCCGCCGGGCATTTAACACCCACCTGGACTTCAGCAGAGTGGTCCGGGGCG AACCAGCCATGGGACGGTCACCCAGCAGTGCCGGGCTGGGCTCTGCGGTGCGGTCCACGGGAGAGC AGGCCTCCAGCTGGAAAGGCCAGGCCTGGAGCTTGCCTCTTCAGTTTTTGTGGCAGTTTTAGTTTT GGGTATTAAAAAGAAAAAAAAAACTTAAAAAAAAAAAAGACACTAACGGCCAGTGAGTTGGAGTCTC AGGGCAGGGTGGCAGTTTCCCTTGAGCAAAGCAGCCAGACGTTGAACTGTGTTTCCTTTCCCTGGG CGCAGGGTGCAGGGTGTCTTCCGGATCTGGTGTGACCTTGGTCCAGGAGTTCTATTTGTTCCTGGG CCTCCTGCGGCATGGGTGTGTCCAGTGCCACCGCTGGCCTCCGCTGCTTCCATCAGCCCTGTCGCC ACCTGGTCCTTCATGAAGAGCAGACACTTAGAGGCTGGTCGGGAATGGGGAGGTCGCCCCTGGGAG GGCAGGCGTTGGTTCCAAGCCGGTTCCCGTCCCTGGCGCCTGGAGTGCACACAGCCCAGTCGGCAC GATCAATCACGTGGACACTAAGGCACGTTTTAGAGTCTCTTGTCTTAATGATTATGTCCATCCGTC TGTCCGTCCATTTGTGTTTTCTGCGTCGTGTCATTGGATATAATCCTCAGAAATAATGCACACTAG CCTCTGACAACCATGAAGCAAAAATCCGTTACATGTGGGTCTGAACTTGTAGACTCGGTCACAGTA TCAAATAAAATCTATAACAGAAAAAAAAAAAAAAAAA

FIGURE 40

MRQTIIKVIKFILIICYTVYYVHNIKFDVDCTVDIESLTGYRTYRCAHPLATLFKILASF
YISLVIFYGLICMYTLWMMLRRSLKKYSFESIREESSYSDIPDVKNDFAFMLHLIDQYDD
LYSKRFAVFLSEVSENKLRQLNLNNEWTLDKLRQRLTKNAQDKLELHLFMLSGIPDTVFD
LVELEVLKLELIPDVTIPPSIAQLTGLKELWLYHTAAKIEAPALAFLRENLRALHIKFTD
IKEIPLWIYSLKTLEELHLTGNLSAENNRYIVIDGLRELKRLKVLRLKSNLSKLPQOVTD
VGVHLQKLSINNEGTKLIVLNSLKKMANLTELELIRCDLERIPHSIFSLHNLQEIDLKDN
NLKTIEEIISFQHLHRLTCLKLWYNHIAYIPTQIGNLTNLERLYLNRNKIEKLPTQLFYC
RKLRYLDLSHNNLTFLPADIGLLQNLQNLAITANRIETLPPELFQCRKLRALHLGNNVLQ
SLPSRVGELTNLTQIELRGNRLECLPVELGECPLLKRSGLVVEEDLFNTLPPEVKERLWR
ADKEOA

Transmembrane domain: amino acids 51-75 (type II)

N-glycosylation site: amino acids 262-266, 290-294, 328-332, 396-400, 432-436, 491-495

cAMP- and cGMP-dependent protein kinase phosphorylation site:

Casein kinase II phosphorylation site: amino acids 91-95, 97-101, 177-181, 253-257, 330-334, 364-368, 398-402, 493-497

N-myristoylation site: amino acids 173-179, 261-267, 395-401, 441-447

FIGURE 41

GGGGGAGAAGGCGGCCGAGCCCCAGCTCTCCGAGCACCGGGTCGGAAGCCGCGACCCGAG AGGTCACCAGCACCCTCGGAACCCAGAGGCCCGCGCTCTGAAGGTGACCCCCCTGGGGAG GAAGGCGATGCCCCTGCGAGGACGATGGCCCGCCCGCCTCGCCCCGGCCGCATCCC GCCCGCGCCCCTGGGCTGCCCGCGGGAGCCGACTGCCTGAACAGCTTTACCGCCGGGGT GCCTGGCTTCGTGCTGGACACCAACGCCTCGGTCAGCAACGGAGCTACCTTCCTGGAGTC CCCACCGTGCGCCGGGGCTGGGACTGCGTGCGCCTGCTGCACCACCAGAACTGCAA CTTGGCGCTAGTGGAGCTGCAGCCCGACCGGGGGAGGACGCCATCGCCGCCTGCTTCCT CAACTACCTCACGAGGGAAGTGTACCGCTCCTACCGCCAGCTGCGGACCCAGGGCTTTGG AGGGTCTGGGATCCCCAAGGCCTGGGCAGGCATAGACTTGAAGGTACAACCCCAGGAACC CCTGGTGCTGAAGGATGTGGAAAACACAGATTGGCGCCTACTGCGGGGTGACACGGATGT CCTGTTCCAGCTGACAGTGACTAGCTCAGACCACCCAGAGGACACGGCCAACGTCACAGT CACTGTGCTGTCCACCAAGCAGACAGAAGACTACTGCCTCGCATCCAACAAGGTGGGTCG CTGCCGGGGCTCTTTCCCACGCTGGTACTATGACCCCACGGAGCAGATCTGCAAGAGTTT CGTTTATGGAGGCTGCTTGGGCAACAAGAACAACTACCTTCGGGAAGAAGAGTGCATTCT AGCCTGTCGGGGTGTGCAAGGTGGGCCTTTGAGAGGCAGCTCTGGGGGCTCAGGCGACTTT CCAGTTCCGCTGCAGCAATGGCTGCTGCATCGACAGTTTCCTGGAGTGTGACGACACCCC CAACTGCCCCGACGCCTCCGACGAGGCTGCCTGTGAAAAATACACGAGTGGCTTTGACGA GCTCCAGCGCATCCATTTCCCCAGTGACAAAGGGCACTGCGTGGACCTGCCAGACACAGG ACTCTGCAAGGAGCATCCCGCGCTGGTACTACAACCCCTTCAGCGAACACTGCGCCCG CTTTACCTATGGTGGTTGTTATGGCAACAAGAACAACTTTGAGGAAGAGCAGCAGTGCCT CGAGTCTTGTCGCGGCATCTCCAAGAAGGATGTGTTTGGCCTGAGGCGGGAAATCCCCAT TCCCAGCACAGGCTCTGTGGAGATGGCTGTCACAGTGTTCCTGGTCATCTGCATTGTGGT GGTGGTAGCCATCTTGGGTTACTGCTTCTTCAAGAACCAGAGAAAGGACTTCCACGGACA CCACCACCACCACCACCCCTGCCAGCTCCACTGTCTCCACTACCGAGGACACGGA GCACCTGGTCTATAACCACACCACCGGCCCCTC<u>TGA</u>GCCTGGGTCTCACCGGCTCTCAC CTGGCCTGCTTCCTGCTTGCCAAGGCAGAGGCCTGGGCTGGGAAAAACTTTGGAACCAG ACTCTTGCCTGTTTCCCAGGCCCACTGTGCCTCAGAGACCAGGGCTCCAGCCCCTCTTGG AGAAGTCTCAGCTAAGCTCACGTCCTGAGAAAGCTCAAAGGTTTGGAAGGAGCAGAAAAC CCTTGGGCCAGAGTACCAGACTAGATGGACCTGCCTGCATAGGAGTTTGGAGGAAGTTG GAGTTTTGTTTCCTCTGTTCAAAGCTGCCTGTCCCTACCCCATGGTGCTAGGAAGAGAG TGGGGTGGTGTCAGACCCTGGAGGCCCCAACCCTGTCCTCCCGAGCTCCTCTTCCATGCT GTGCGCCCAGGGCTGGGAGGAAGGACTTCCCTGTGTAGTTTGTGCTGTAAAGAGTTGCTT TTTGTTTATTTAATGCTGTGGCATGGGTGAAGAGGGGGGAAGAGGCCTGTTTGGCCTCT CTGTCCTCTCTCTCTCCCCCAAGATTGAGCTCTCTGCCCTTGATCAGCCCCACCCTG GCCTAGACCAGCAGACAGAGCCAGGAGAGGCTCAGCTGCATTCCGCAGCCCCCACCCCCA AGGTTCTCCAACATCACAGCCCAGCCCACCCACTGGGTAATAAAAGTGGTTTGTGGAAAA AAAAAAAAAAAAAAAAAAA

FIGURE 42

MAPARTMARARLAPAGIPAVALWLLCTLGLQGTQAGPPPAPPGLPAGADCLNSFTAGVPG
FVLDTNASVSMGATFLESPTVRRGWDCVRACCTTQNCNLALVELQPDRGEDAIAACFLIN
CLYEQNFVCKFAPREGFINYLTREVYRSYRQULTQGFGGGIPKAWAGIDLKVQPQEBPLV
LKDVENTDWRLLRGDTDVRVERKDPNQVELWGLKEGTYLFQLTVTSSDHPEDTANVTVTV
LSTKQTEDYCLASNKVGRCRGSFFRWYNDFTEGICKSFVYGGCLGNKNNYLREEECILAC
RGVQGGPLRGSSGAQATFFQGSBMERRHPVCSGTCQPTQFRCSNGCCIDSFLECDDTPNC
PDASDEAACEKYTSGFDELQRIHFPSDKGHCVDLPDTGLCKESIPRWYYNPFSEHCARFT
YGGCYGNKNNFEEEQQCLESCRGISKKDVFGLRREIPIPSTGSVEMAVTVFLVICIVVV
AILGYCFFKNQRKDFHGHHHHPPPTPASSTVSTTEDTEHLVYNHTTRPL

signal sequence: Amino acids 1-35

transmembrane domain: Amino acids 466-483

N-glycosylation sites: Amino acids 66-70;235-239;523-527

N-myristoylation sites:

A m i n o a c i d s 29-35;43-49;161-167;212-218;281-287;282-288;285-291; 310-316;313-319;422-428;423-429;426-432

Cell attachment sequence: Amino acids 193-199

Pancreatic trypsin inhibitor (Kunitz) family signatures: Amino acids 278-298;419-438

FIGURE 43

CCCACGCGTCCGCACCTCGGCCCCGGGCTCCGAAGCGGCTCGGGGGCGCCCTTTCGGTCA ACATCGTAGTCCACCCCTCCCCATCCCCAGCCCCCGGGGATTCAGGCTCGCCAGCGCCC AGC/AGGGAGCCGGCAGGGAAGCGCGATGGGGGCCCCAGCCGCCTCGCTCCTGCT TCCTGCTGTTCGCCTGCTGGCGGCGCCGGCGGGGCCAACCTCTCCCAGGACGACACACC AGCCCTGGACATCTGATGAAACAGTGGTGGCTGGTGGCACCGTGGTGCTCAAGTGCCAAG TGAAAGATCACGAGGACTCATCCCTGCAATGGTCTAACCCTGCTCAGCAGACTCTCTACT TTGGGGAGAGAGAGCCCTTCGAGATAATCGAATTCAGCTGGTTACCTCTACGCCCCACG AGCTCAGCATCAGCATCAGCAATGTGGCCCTGGCAGACGAGGGCGAGTACACCTGCTCAA TCTTCACTATGCCTGTGCGAACTGCCAAGTCCCTCGTCACTGTGCTAGGAATTCCACAGA AGCCCATCATCACTGGTTATAAATCTTCATTACGGGAAAAAGACACAGCCACCCTAAACT GTCAGTCTTCTGGGAGCAAGCCTGCAGCCCGGCTCACCTGGAGAAAGGGTGACCAAGAAC TCCACGGAGAACCAACCGCATACAGGAAGATCCCAATGGTAAAACCTTCACTGTCAGCA GCTCGGTGACATTCCAGGTTACCCGGGAGGATGATGGGGGCGAGCATCGTGTGCTCTGTGA ACCATGAATCTCTAAAGGGAGCTGACAGATCCACCTCTCAACGCATTGAAGTTTTATACA CACCAACTGCGATGATTAGGCCAGACCCTCCCCATCCTCGTGAGGGCCAGAAGCTGTTGC TACACTGTGAGGGTCGCGGCAATCCAGTCCCCCAGCAGTACCTATGGGAGAAGGAGGGCA GTGTGCCACCCTGAAGATGACCCAGGAGAGTGCCCTGATCTTCCCTTTCCTCAACAAGA GTGACAGTGGCACCTACGGCTGCACAGCCACCAGCAACATGGGCAGCTACAAGGCCTACT ACACCCTCAATGTTAATGACCCCAGTCCGGTGCCCTCCTCCTCCAGCACCTACCACGCCA TCATCGGTGGGATCGTGGCTTTCATTGTCTTCCTGCTGCTCATCATGCTCATCTTCCTTG GCCACTACTTGATCCGGCACAAAGGAACCTACCTGACACATGAGGCAAAAAGGCTCCGACG ATGCTCCAGACGCGGACACGGCCATCATCAATGCAGAAGGCGGGCAGTCAGGAGGGGACG ACAAGAAGGAATATTTCATC<u>TAG</u>AGGCGCCTGCCCACTTCCTGCGCCCCCCAGGGGCCCT GTGGGGACTGCTGGGGCCGTCACCAACCCGGACTTGTACAGAGCAACCGCAGGGCCGCCC CTCCCGCTTGCTCCCCAGCCCACCCACCCCCTGTACAGAATGTCTGCTTTGGGTGCGGT CCCTTTCCGTGGCTTCTCTGCATTTGGGTTATTATTATTTTTTGTAACAATCCCAAATCAA ATCTGTCTCCAGGCTGGAGAGGCAGGAGCCCTGGGGTGAGAAAAAGCAAAAAACAAA AAACA

FIGURE 44

MGAPAASLLLLLLLFACCWAPGGANLSQDDSQPWTSDETVVAGGTVVLKCQVKDHEDSSL QWSMPAQQTLYFGEKRALRDNRIQLVTSTPHELSISISNVALADEGEYTCSIFTMEVRTI KSLVTVLGIPQKPIITGYKSSLREKDTATLMCQSSGSKPAARLTWRGDQELHGGEPTRIQ EDPNGKTFTVSSSVTFQVTREDDGASIVCSVNHESLKGADRSTSQRIEVLYTPTAMIRPD PPHPREGQKLLLHCEGRCNPVPQQYLWEKEGSVPPLKMTQESALIFFFLNKSDSGTYGCT ATSNMGSYKAYYTLNVNDPSPVPSSSSTYHAIIGGIVAFIVFLLLIMLIFLGHYLIRHKG TYLTHEAKGSDDAPDADTAIINAEGGQSGGDKKEYFI

Signal sequence:

Transmembrane domain: amino acids 331-352

N-glycosylation site: amino acids 25-29, 290-294

Casein kinase II phosphorylation site: amino acids 27-31, 35-39, 89-93, 141-145, 199-203, 388-392

N-myristoylation site: amino acids 2-8, 23-29, 156-162, 218-224, 295-301, 298-304, 306-310, 334-340, 360-364, 385-389, 386-390

Prokaryotic membrane lipoprotein lipid attachment site: amino acids 7-18

FIGURE 45

ACTTGCCATCACCTGTTGCCAGTGTGGAAAAATTCTCCCTGTTGAATTTTTTTGCACATGG AGGACAGCAGCAAAGAGGGCAACACAGGCTGATAAGACCAGAGACAGCAGGGAGATTATT TTACCATACGCCCTCAGGACGTTCCCTCTAGCTGGAGTTCTGGACTTCAACAGAACCCCA TATTTATTTCCGTACTTCAGAAATGGGCCTACAGACCACAAAGTGGCCCAGCCATGGGG CTTTTTTCCTGA AGTCTTGGCTTATCATTTCCCTGGGGCTCTACTCACAGGTGTCCAAAC TCCTGGCCTGCCCTAGTGTGTGCCGCTGCGACAGGAACTTTGTCTACTGTAATGAGCGAA GCTTGACCTCAGTGCCTCTTGGGATCCCGGAGGGCGTAACCGTACTCTACCTCCACAACA ACCAAATTAATAATGCTGGATTTCCTGCAGAACTGCACAATGTACAGTCGGTGCACACGG TCTACCTGTATGGCAACCAACTGGACGAATTCCCCATGAACCTTCCCAAGAATGTCAGAG TTCTCCATTTGCAGGAAAACAATATTCAGACCATTTCACGGGCTGCTCTTGCCCAGCTCT TGAAGCTTGAAGAGCTGCACCTGGATGACAACTCCATATCCACAGTGGGGGTGGAAGACG GGGCCTTCCGGGAGGCTATTAGCCTCAAATTGTTGTTTTTTGTCTAAGAATCACCTGAGCA GTGTGCCTGTTGGGCTTCCTGTGGACTTGCAAGAGCTGAGAGTGGATGAAAATCGAATTG CTGTCATATCCGACATGGCCTTCCAGAATCTCACGAGCTTGGAGCGTCTTATTGTGGACG GGAACCTCCTGACCAACAAGGGTATCGCCGAGGGCACCTTCAGCCATCTCACCAAGCTCA AGGA ATTTTCA ATTGTACGTA ATTCGCTGTCCCACCCTCCTCCCGATCTCCCAGGTACGC ATCTGATCAGGCTCTATTTGCAGGACAACCAGATAAACCACATTCCTTTGACAGCCTTCT AAGGGGTTTTTGATAATCTCTCCAACCTGAAGCAGCTCACTGCTCGGAATAACCCTTGGT TTTGTGACTGCAGTATTAAATGGGTCACAGAATGGCTCAAATATATCCCTTCATCTCTCA ACGTGCGGGGTTTCATGTGCCAAGGTCCTGAACAAGTCCGGGGGATGGCCGTCAGGGAAT CCCCAAGTACAGCTTCTCCGACCACTCAGCCTCCCACCCTCTCTATTCCAAACCCTAGCA GAAGCTACACGCCTCCAACTCCTACCACATCGAAACTTCCCACGATTCCTGACTGGGATG GCAGAGAAAGAGTGACCCCACCTATTTCTGAACGGATCCAGCTCTCTATCCATTTTGTGA ATGATACTTCCATTCAAGTCAGCTGGCTCTCTCTCTCTCACCGTGATGGCATACAAACTCA CATGGGTGAAAATGGGCCACAGTTTAGTAGGGGGGCATCGTTCAGGAGCGCATAGTCAGCG GTGAGAACCACCTGAGCCTGGTTAACTTAGAGCCCCGATCCACCTATCGGATTTGTT TAGTGCCACTGGATGCTTTTAACTACCGCGCGGTAGAAGACACCATTTGTTCAGAGGCCA $\tt CGTCCCACAGCATGGGCTCCCCCTTTCTGCTGGCGGGCTTGATCGGGGGCGCGGTGATAT$ TTGTGCTGGTGTCTTGCTCAGCGTCTTTTGCTGGCATATGCACAAAAAGGGGCGCTACA CCAAGAAGGACAACTCCATCCTGGAGATGACAGAAACCAGTTTTCAGATCGTCTCCTTAA ATAACGATCAACTCCTTAAAGGAGATTTCAGACTGCAGCCCATTTACACCCCAAATGGGG GCATTAATTACACAGACTGCCATATCCCCAACAACATGCGATACTGCAACAGCAGCGTGC CAGACCTGGAGCACTGCCATACGTGACAGCCAGAGGCCCAGCGTTATCAAGGCGGACAAT TAGACTCTTGAGAACACACTCGTGTGTGCACATAAAGACACGCAGATTACATTTGATAAA TGTTACACAGATGCATTTGTGCATTTGAATACTCTGTAATTTATACGGTGTACTATATAA ${\tt TGGGATTTAAAAAAAGTGCTATCTTTTCTATTTCAAGTTAATTACAAACAGTTTTGTAAC}$ TCTTTGCTTTTTAAATCTT

FIGURE 46

MGLQTTKWPSHGAFFLKSWLIISLGLYSQVSKLLACPSVCRCDRNFVYCNERSLTSVPLG
IPEGVTVLYLHNNQINNAGFPAELHNVQSVHTVYLYGNQLDEFPMNLPKNVRVLHLQENN
IQTISRAALAQLLKLEELHLDDNSISTVGVEEGAFREAISLKLLFLSKNHLSSVPVGLPV
DLQBLRVDENRIAVISDMAFQNLTSLERLIVDGNLLTNKGIABGTFSHLTKLKEFSIVRN
SLSHPPPDLPGTHLIRLYLQDNQINHIPLTAFSNLRKLERLDISNNQLRMLTGGVFDNLS
NLKQLTARNNPWFCDCSIKWVTEWLKYIPSSLNVRGFMCQGPEQVRGMAVRELNMNLLSC
PTTTPGLPLFTPAPSTASPTTQPPTLSIPNPSRSYTPPTTTSKLPTIPDMDGRERVTPF
ISERIQLSIHFVMDTSIQVSWLSLFTVMAYKLTWVKMGHSLVGGIVQERIVSGEKQHLSL
VNLEPRSTYRICLVPLDAFNYRAVEDTICSEATTHASYLNNGSNTASSHEQTTSHSMGSP
FLLAGLIGGAVIFVLVVLLSVFCWHMHKKGRYTYSQKWKYNRGRKDDYCEAGTKKDNSI
EMTETSFQIVSLNNDQDLLKGDPRLQPIYTPNGGINYTDCHIPNNMRYCNSSVPDLEHCHT

Signal peptide:

Transmembrane domain:

N-glycosylation site: amino acids 202-206, 298-302, 433-437, 521-525, 635-639, 649-653

Casein kinase II phosphorylation site: amino acids 204-208, 407-411, 527-531, 593-597, 598-602, 651-655

Tyrosine kinase phosphorylation site: amino acids 319-328

N-myristoylation site: amino acids 2-8, 60-66, 149-155, 213-219, 220-226, 294-300, 522-528, 545-551, 633-639

Amidation site: amino acids 581-585

Leucine zipper pattern: amino acids 164-186

Phospholipase A2 aspartic acid active site: amino acids 39-50

FIGURE 47

GCAGCGAGCGCCGGGTGCGGCCCTGCCGCCGCAGGGATGTGACCTTCACCGTCGCTTAGC CAGGATGACCGGAGCCCGTGTCTCGCGGCGTCCGCGCCTCGCTTCAGCCTCCCGGGTGCT CCGGCCCGCCGCCCTGGGCACTGCGCTTCTGCTGCTCCTGCTGCTTCCGAGTCTT CTCACACTGTGCTGTTGCGGGCGCGTGAGGCGGCGCAGTTTCTGCGGCCCAGGCAGCCC GCGCCTACCAAGTCTTCGAGGAGGCCAAGCAGGCCCACCTGGAACGGGAGTGCGTGGAGG AGGTGTGCAGCAAAGAGGGGCCAGAGAGGTGTTCGAGAACGACCCCGAGACGGAGTATT TCTATCCACGATATCAAGAGTGCATGAGAAAATATGGCAGGCCTGAAGAAAAAAACCCAG ATTTCGCCAAATGTGTTCAGAACTTGCCTGACCAGTGCACCCCAAACCCTTGTGATAAGA AGGGTACTCATATCTGCCAAGACCTCATGGGCAACTTCTTCTGCGTGTGCACAGATGGCT GGGGAGGCCGCTCTGTGACAAAGATGTCAATGAGTGTGTCCAGAAGAATGGGGGCTGCA TTGCATCAGACGGCCAGACCTGCCAAGATATCGATGAATGCACAGACTCAGACACCTGTG GGGACGCGCGATGCAAGAACTTGCCAGGCTCCTACTCTTGCCTCTGCGATGAGGGATATA CATACAGCTCCAAGGAGAGACCTGCCAAGATGTGGACGAGTGCCAGCAGGATCGCTGTG ACCAGA CCTGTGTCA ACTCCCCAGGCAGCTATACCTGCCACTGTGATGGGCGAGGGGGCC TAAAACTATCCCCAGACATGGATACTTGTGAGGACATCTTACCATGTGTGCCCTTCAGCA TGGCCAAGAGCGTGAAGTCCTTGTACCTGGGCCGCATGTTCAGCGGGACCCCCGTGATTA TTGACCCTGAAGGAGTCCTCTTCTTCGCTGGAGGCCGTTCAGACAGCACCTGGATTGTCC TGGGCCTAAGAGCTGGGCGGCTTGAGCTGCAGCTTCGGTACAATGGCGTTGGGCGCATCA CCAGCAGCGGGCCAACCATCAACCACGGCATGTGGCAAACTATCTCCGTGGAAGAGCTGG AACGTAACCTTGTCATCAAGGTCAACAAAGATGCTGTAATGAAGATCGCGGTAGCTGGGG AGCTGTTTCAGCTGGAGAGGGGCCTCTATCACCTGAATCTCACCGTGGGCGGCATTCCCT TCAAGGAGAGTGAGCTCGTCCAGCCGATTAACCCTCGCCTGGATGGGTGCATGAGGAGTT GGAACTGGCTGAACGGGGAAGACAGCGCCATCCAGGAGACAGTCAAGGCAAACACAAAAA TGCAGTGCTTCTCTGTGACAGAAAGGGGCTCCTTCTTCCCGGGGAATGGATTTGCTACCT ACAGGCTCAACTACACCCGAACATCGCTGGATGTCGGCACGGAAACCACCTGGGAAGTTA AAGTTGTGGCTCGGATCCGCCCTGCCACGGACACGGGGGTGCTGCTGGCGCTGGTGGGGG ACGACGATGTCGTCATCTCTGTGGCCCTAGTCGACTACCACTCTACAAAGAAGCTCAAGA AGCAGTTGGTGGTCCTGGCAGTTGAGGATGTTGCCCTGGCACTGATGGAAATCAAGGTGT GCGACAGCCAGGAACACACGGTCACTGTCTCCCTGCGGGAGGGTGAGGCCACCCTAGAAG TGGATGGCACAAAGGGCCAGAGTGAAGTGAGCACTGCCCAGCTGCAGGAGCGACTGGACA CACTTAAGACACATCTGCAAGGCTCTGTGCACACCTATGTTGGAGGCCTGCCAGAAGTAT CGGTGATTTCTGCACCCGTCACTGCGTTCTACCGCGGATGCATGACTCTGGAGGTAAACG GGAAAATCCTGGACCTGGATACGGCCTCGTACAAGCACAGTGACATCACCTCCCACTCCT GCCGGCTGTGGAGCATGCCACCCCTAGACCGAGCTGCAAGAGGGCTCCACACCTAAAG ACAAAAATGAAGCAGGGTTTGGACACACAGCACTGGCTCCTCTCGCATGGTCCTGCAACA GACCTGGTGGGAGCTGGCTGGAAGGGGCTGGGCTAGAGGGGCTGGCAGTTTGCAGCAGAA

FIGURE 48

MPPPPGPAAALGTALLLLLASESSHTVLLRAREAAQFLRPRQRRAYQVFEEAKQGHLER ECVEBVCSKEBAREVFENDPETEYFYPRYDECMRKYGRPEEKNPDFAKCVQNLPDQCTPN PCDKKGTHICQDLMGNFFCVCTDGWGGRLCDKDVNECVQKNGGCSQVCHNKPGSFQCACH SGFSLASDGQTCQDIDECTDSDTCGDARCKNLPGSYSCLCDEGYTYSSKEKTCQDVDECQ QDRCEQTCVNSPGSYTCHCDGRGGLKLSPDMDTCEDILPCVPFSMAKSVKSLYLGRMFSG TPVIRLFRKRLQPTRLLAEFDFFTFDPESVLFFAGGRSDSTNIVLGLRAGRLELQLRYNG VGRITSSGPTINHGMWQTISVEELERNLVIKVNKDAVMKIAVAGELFCLERGLYHLNLTV GGIPFKESELVQPINPRLDGCMRSWNWLNGEDSAIQETVKANTKMQCFSVTERGSFFPGN GFATYRLNYTRTSLDVGTETTWEVKVVARIRPATDTGVLLALVGDDDVVISVALVDYHST KKLKKQLVVLAVEDDALALMEIKVCDSQEHTVTVSLREGEATLEVDGTKGQSEVSTAQLQ ERLDTLKTHLQGSVHTYVGGLPEVSISAPVTAFYRGCMTLEVNGKILDLDTASYKHSDI TSHSCPPVEHATP

FIGURE 49

CCGGCGGCCTCCCGGCGGAGCGAGCAGATCCAGTCCGGCCCGCAGCGCAACTCGGTCCA $\tt GTCGGGGCGGCGCTGCGGGCGCAGAGCGGAGATGCAGCGGCTTGGGGCCACCCTGCTGT$ GCCTGCTGCTGGCGGCGGCGGTCCCCACGGCCCCGCGCCCGCTCCGACGGCGACCTCGG CTCCAGTCAAGCCCGGCCCGGCTCTCAGCTACCCGCAGGAGGAGGCCACCCTCAATGAGA TGTTCCGCGAGGTTGAGGAACTGATGGAGGACACGCAGCACAAATTGCGCAGCGCGGTGG A AGAGA TGGA GGCAGA AGA AGCTGCTGCTA AAGCATCATCAGA AGTGAACCTGGCAAACT TACCTCCCAGCTATCACAATGAGACCAACACAGACACGAAGGTTGGAAATAATACCATCC ATGTGCACCGAGAAATTCACAAGATAACCAACAACCAGACTGGACAAATGGTCTTTTCAG AGACAGTTATCACATCTGTGGGAGACGAAGAAGGCAGAAGGAGCCACGAGTGCATCATCG ACGAGGACTGTGGGCCCAGCATGTACTGCCAGTTTGCCAGCTTCCAGTACACCTGCCAGC GTGTCTGGGGTCACTGCACCAAAATGGCCACCAGGGGCAGCAATGGGACCATCTGTGACA ACCAGAGGGACTGCCAGCCGGGGCTGTGCTGTGCCTTCCAGAGAGGCCTGCTGTTCCCTG TGTGCACACCCCTGCCCGTGGAGGGCGAGCTTTGCCATGACCCCGCCAGCCGGCTTCTGG ACCTCATCACCTGGGAGCTAGAGCCTGATGGAGCCTTGGACCGATGCCCTTGTGCCAGTG GCCTCCTCTGCCAGCCCCACAGCCACAGCCTGGTGTATGTGTGCAAGCCGACCTTCGTGG GGAGCCGTGACCAAGATGGGGAGATCCTGCTGCCCAGAGAGGTCCCCGATGAGTATGAAG AAGAGATGGCGCTGGGGGAGCCTGCGGCTGCCGCCGCTGCACTGCTGGGAGGGGAAGAGA GCTGTGTGTGTTTTAGGCGTGGGCTGACCAGGCTTCTTCCTACATCTTCCTACAGTTAAGTT TCCCCTCTGGCTTGACAGCATGAGGTGTTGTGCATTTGTTCAGCTCCCCCAGGCTGTTCT CCAGGCTTCACAGTCTGGTGCTTGGGAGAGTCAGGCAGGGTTAAACTGCAGGAGCAGTTT GCCACCCTGTCCAGATTATTGGCTGCTTTGCCTCTACCAGTTGGCAGACAGCCGTTTGT TCTACATGGCTTTGATAATTGTTTGAGGGGAGGAGATGGAAACAATGTGGAGTCTCCCTC TGATTGGTTTTGGGGAAATGTGGAGAAGAGTGCCCTGCTTTGCAAACATCAACCTGGCAA AAATGCAACAAATGAATTTTCCACGCAGTTCTTTCCATGGGCATAGGTAAGCTGTGCCTT CAGCTGTTGCAGATGAAATGTTCTGTTCACCCTGCATTACATGTGTTTATTCATCCAGCA GTGTTGCTCAGCTCCTACCTCTGTGCCAGGGCAGCATTTTCATATCCAAGATCAATTCCC TCTCTCAGCACAGCCTGGGGAGGGGGTCATTGTTCTCCTCGTCCATCAGGGATCTCAGAG GCTCAGAGACTGCAAGCTGCTTGCCCAAGTCACACAGCTAGTGAAGACCAGAGCAGTTTC ATCTGGTTGTGACTCTAAGCTCAGTGCTCTCTCCACTACCCCACACCAGCCTTGGTGCCA CCAAAAGTGCTCCCCAAAAGGAAGGAGAATGGGATTTTTCTTGAGGCATGCACATCTGGA ATTAAGGTCAAACTAATTCTCACATCCCTCTAAAAGTAAACTACTGTTAGGAACAGCAGT GTTCTCACAGTGTGGGGCAGCCGTCCTTCTAATGAAGACAATGATATTGACACTGTCCCT CTTTGGCAGTTGCATTAGTAACTTTGAAAGGTATATGACTGAGCGTAGCATACAGGTTAA CCTGCAGAAACAGTACTTAGGTAATTGTAGGGCGAGGATTATAAATGAAATTTGCAAAAT CACTTAGCAGCAACTGAAGACAATTATCAACCACGTGGAGAAAATCAAACCGAGCAGGGC TGTGTGAAACATGGTTGTAATATGCGACTGCGAACACTGAACTCTACGCCACTCCACAAA TGATGTTTTCAGGTGTCATGGACTGTTGCCACCATGTATTCATCCAGAGTTCTTAAAGTT TAAAGTTGCACATGATTGTATAAGCATGCTTTCTTTGAGTTTTAAATTATGTATAAACAT AAAAAA

FIGURE 50

MQRLGATLLCLLLAAAVPTAPAPAPTATSAPVKPGPALSYPQEEATLNEMFREVEELMED
TQHKLRSAVEEMBAEBAAKASSEVNLANLPPSYHBETNITDIKVGNNTIHVHREIHKITN
NQTGGMVFSETVITSVGDEEGRSHECIIDEDCGPSMYCQFASFQYTCQPCRGQMLCTR
DSECCGDQLCVWGHCTKMATRGSNGTICDNQRDCQPGLCCAFQRGLLFPVCTPLPVEGEL
CHDPASRLLDLITWELEPDGALDRCPCASGLLCQPHSHSLVYVCKPTFVGSRDQDGEILL
PREVPDEYPVGSFMEEVROELEDLERSITEEMALGEPAAAAAALLGGEEI

Signal sequence: amino acids 1-19

N-glycosylation site: amino acids 96-100, 106-110, 121-125, 204-208

Casein kinase II phosphorylation site: amino acids 46-50, 67-71, 98-102, 135-139, 206-210, 312-316, 327-331

N-myristoylation site: amino acids 202-208, 217-223

Amidation site: amino acids 140-144

FIGURE 51

GCCTGTTGCTGATGCTGCCGTGCGGTACTTGTCATGGAGCTGGCACTGCGGCGCTCTCCC GACTGGCCCACAGAGGAGGGCAAGGAAGTATGGGATTATGTGACGGTCCGCAAGGATGCC TACATGTTCTGGTGGCTCTATTATGCCACCAACTCCTGCAAGAACTTCTCAGAACTGCCC CTGGTCATGTGGCTTCAGGGCGGTCCAGGCGGTTCTAGCACTGGATTTGGAAACTTTGAG GAAATTGGGCCCCTTGACAGTGATCTCAAACCACGGAAAACCACCTGGCTCCAGGCTGCC AGTCTCCTATTTGTGGATAATCCCGTGGGCACTGGGTTCAGTTATGTGAATGGTAGTGGT GCCTATGCCAAGGACCTGGCTATGGTGGCTTCAGACATGATGGTTCTCCTGAAGACCTTC TTCAGTTGCCACAAAGAATTCCAGACAGTTCCATTCTACATTTTCTCAGAGTCCTATGGA GGAAAAATGGCAGCTGGCATTGGTCTAGAGCTTTATAAGGCCATTCAGCGAGGGACCATC AAGTGCAACTTTGCGGGGGTTGCCTTGGGTGATTCCTGGATCTCCCCTGTTGATTCGGTG CTCTCCTGGGGACCTTACCTGTACAGCATGTCTCTCTCGAAGACAAAGGTCTGGCAGAG GTGTCTAAGGTTGCAGAGCAAGTACTGAATGCCGTAAATAAGGGGCTCTACAGAGAGGCC ACAGAGCTGTGGGGGAAAGCAGAAATGATCATTGAACAGAACACAGATGGGGTGAACTTC TATAACATCTTAACTAAAAGCACTCCCACGTCTACAATGGAGTCGAGTCTAGAATTCACA CAGAGCCACCTAGTTTGTCTTTGTCAGCGCCACGTGAGACACCTACAACGAGATGCCTTA AGCCAGCTCATGATGGCCCCATCAGAAAGAAGCTCAAAATTATTCCTGAGGATCAATCC TGGGGAGGCCAGGCTACCAACGTCTTTGTGAACATGGAGGAGGACTTCATGAAGCCAGTC ATTAGCATTGTGGACGAGTTGCTGGAGGCAGGGATCAACGTGACGGTGTATAATGGACAG CTGGATCTCATCGTAGATACCATGGGTCAGGAGGCCTGGGTGCGGAAACTGAAGTGGCCA GAACTGCCTAAATTCAGTCAGCTGAAGTGGAAGGCCCTGTACAGTGACCCTAAATCTTTG GAAACATCTGCTTTTGTCAAGTCCTACAAGAACCTTGCTTTCTACTGGATTCTGAAAGCT GGTCATATGGTTCCTTCTGACCAAGGGGACATGGCTCTGAAGATGATGAGACTGGTGACT CAGCAAGAATAGGATGGATGGGGCTGGAGATGAGCTGGTTTGGCCTTGGGGCACAGAGCT GAGCTGAGGCCGCTGAAGCTGTAGGAAGCGCCATTCTTCCCTGTATCTAACTGGGGCTGT CATCA AGA AGGTTCTGACCAGCTTCTGCAGAGGATAAAATCATTGTCTCTGGAGGCAATT TGGAAATTATTTCTGCTTCTTAAAAAAACCTAAGATTTTTTAAAAAAATTGATTTTTT ATCAAAATAAAGGATGATAATAGATATTAA

FIGURE 52

MELALRRSPVPRNLLLLELLGLNAGAVIDWPTEEGKEVWDVVTVRKDAYMFWWLLYYATN SCKNFSELPLVMWLQGGPGGSSTGFGNFEEIGPLDSDLKPRKTTWLQAASLLFVDNPVGT GFSYVNGSGAYAKDLAMVASDMMVLLKTFFSCHKEFQTVPFYIFSESYGGKMAAGIGLEL YKAIQRGTIKCNFAGVALGDSWISPVDSVLSWGPYLYSMSLLEDKGLAEVSKVAEQUIND VNKGLYREATELWGKAEMIIEQNTDGVNFFYNILTKSTPFSTMESSLEFTQSHLVCLCQRH VRHLQRDALSQLMNGPIRKKLKIIPEDQSWGGQATNVFVNMEEDFMKPVISIVDELLEAG INTYVYNGGLDLIVDTMGQEAWVRKLKWPELPKFSQLKWKALYSDPKSLETSAFVKSYKN LAFYWILKAGHMVPSDOGDMALKMMRLVTOOE

Signal sequence: amino acids 1-25

N-glycosylation site: amino acids 64-68, 126-130, 362-366

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 101-105

Casein kinase II phosphorylation site: amino acids 204-208, 220-224, 280-284, 284-288, 351-355, 449-453

N-myristoylation site: amino acids 22-28, 76-82, 79-85, 80-86, 119-125, 169-175, 187-193, 195-201, 331-337, 332-338, 360-366

FIGURE 53

GTCTGTTCCCAGGAGTCCTTCGGCGGCTGTTGTGTCAGTGGCCTGATCGCGATGGGGACA AAGGCGCAAGTCGAGAGGAAACTGTTGTGCCTCTTCATATTGGCGATCCTGTTGTGCTCC CTGGCATTGGGCAGTGTTACAGTGCACTCTTCTGAACCTGAAGTCAGAATTCCTGAGAAT AATCCTGTGAAGTTGTCCTGTGCCTACTCGGGCTTTTCTTCTCCCCGTGTGGAGTGGAAG TTTGACCAAGGAGACACCACCAGACTCGTTTGCTATAATAACAAGATCACAGCTTCCTAT GAGGACCGGGTGACCTTCTTGCCAACTGGTATCACCTTCAAGTCCGTGACACGGGAAGAC ACTGGGACATACACTTGTATGGTCTCTGAGGAAGGCGGCAACAGCTATGGGGAGGTCAAG GTCAAGCTCATCGTGCTTGTGCCTCCATCCAAGCCTACAGTTAACATCCCCTCCTCTGCC ACCATTGGGAACCGGGCAGTGCTGACATGCTCAGAACAAGATGGTTCCCCACCTTCTGAA TACACCTGGTTCAAAGATGGGATAGTGATGCCTACGAATCCCAAAAGCACCCGTGCCTTC AGCAACTCTTCCTATGTCCTGAATCCCACAACAGGAGAGCTGGTCTTTGATCCCCTGTCA GCCTCTGATACTGGAGAATACAGCTGTGAGGCACGGAATGGGTATGGGACACCCATGACT TCAAATGCTGTGCGCATGGAAGCTGTGGAGCGGAATGTGGGGGTCATCGTGGCAGCCGTC CTTGTAACCCTGATTCTCCTGGGAATCTTGGTTTTTGGCATCTGGTTTGCCTATAGCCGA AGTGCCCGAAGTGAAGGAGAATTCAAACAGACCTCGTCATTCCTGGTGTGAGCCTGGTCG GCTCACCGCCTATCATCTGCATTTGCCTTACTCAGGTGCTACCGGACTCTGGCCCCTGAT GTCTGTAGTTTCACAGGATGCCTTATTTGTCTTCTACACCCCACAGGGCCCCCTACTTCT TTTCCTACCACTGCTGAGTGGCCTGGAACTTGTTTAAAGTGTTTATTCCCCATTTCTTTG AGGGATCAGGAAGGAATCCTGGGTATGCCATTGACTTCCCTTCTAAGTAGACAGCAAAAA AGGTATCTTGAGCTTGGTTCTGGGCTCTTTCCTTGTGTACTGACGACCAGGGCCAGCTGT TCTAGAGCGGGAATTAGAGGCTAGAGCGGCTGAAATGGTTGTTTGGTGATGACACTGGGG TCCTTCCATCTGGGGCCCACTCTCTTCTGTCTTCCCATGGGAAGTGCCACTGGGATCC AGCTCTTGTTGTGGAGAGCATAGTAAATTTTCAGAGAACTTGAAGCCAAAAGGATTTAAA ACCGCTGCTCTAAAGAAAAGAAAACTGGAGGCTGGGCGCAGTGGCTCACGCCTGTAATCC CAGAGGCTGAGGCAGGCGGATCACCTGAGGTCGGGAGTTCGGGATCAGCCTGACCAACAT GGAGAAACCCTACTGGAAATACAAAGTTAGCCAGGCATGGTGGTGCATGCCTGTAGTCCC

FIGURE 54

MGTKAQVERKLLCLFILAILLCSLALGSVTVHSSEPEVRIPENNPVKLSCAYSGFSSPRV EWKFDQGDTTRLVCYNNKITASYEDRVTFLPTGITFKSVTREDTGTYTCMVSEEGGNSYG EVKVKLIVLVPPSKPTVNIPSSATIGNRAVLTCSEQDGSPPSEYTWFKDGIVMPTNPKST RAFSNSSYVLNPTTGELVFDPLSASDTGEYSCEARNGYGTPMTSNAVRMEAVERNVGVIV AAVLVTLILLGILVFGIWFAYSRGHFDRTKKGTSSKKVIYSQPSARSEGEFKQTSSFLV

Signal sequence: amino acids 1-27

Transmembrane domain: amino acids 238-255

N-glycosylation site: amino acids 185-189

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 270-274

Casein kinase II phosphorylation site: amino acids 34-38, 82-86, 100-104, 118-122, 152-156, 154-158, 193-197, 203-207, 287-291

N-myristoylation site: amino acids 105-111, 116-122, 158-164, 219-225, 237-243, 256-262

FIGURE 55

GTTGTGTCCTTCAGCAAAACAGTGGATTTAAATCTCCTTGCACAAGCTTGAGAGCAACAC AAGAAAAAAATCATGAAAACCATCCAGCCAAAAATGCACAATTCTATCTCTTGGGCAAT CTTCACGGGGCTGCTCTCTGTGTCTCTTCCAAGGAGTGCCCGTGCGCAGCGGAGATGC CACCTTCCCCAAAGCTATGGACAACGTGACGGTCCGGCAGGGGGAGAGCGCCACCCTCAG GTGCACTATTGACAACCGGGTCACCCGGGTGGCCTGGCTAAACCGCAGCACCATCCTCTA TGCTGGGAATGACAAGTGGTGCCTGGATCCTCGCGTGGTCCTTCTGAGCAACACCCAAAC GCAGTACAGCATCGAGATCCAGAACGTGGATGTGTATGACGAGGGCCCTTACACCTGCTC GGTGCAGACAGCCACCCAAAGACCTCTAGGGTCCACCTCATTGTGCAAGTATCTCC CAAAATTGTAGAGATTTCTTCAGATATCTCCATTAATGAAGGGAACAATATTAGCCTCAC CTGCATAGCAACTGGTAGACCAGAGCCTACGGTTACTTGGAGACACATCTCTCCCAAAGC GGTTGGCTTTGTGAGTGAAGACGAATACTTGGAAATTCAGGGCATCACCCGGGAGCAGTC AGGGGACTACGAGTGCAGTGCCTCCAATGACGTGGCCGCGCCCGTGGTACGGAGAGTAAA GGTCACCGTGAACTATCCACCATACATTTCAGAAGCCAAGGGTACAGGTGTCCCCGTGGG ACAAAAGGGGACACTGCAGTGTGAAGCCTCAGCAGTCCCCTCAGCAGAATTCCAGTGGTA CAAGGATGACAAAAGACTGATTGAAGGAAAGAAAGGGGTGAAAGTGGAAAACAGACCTTT CCTCTCAAAACTCATCTTCTTCAATGTCTCTGAACATGACTATGGGAACTACACTTGCGT GGCCTCCAACAAGCTGGGCCACACCAATGCCAGCATCATGCTATTTGGTCCAGGCGCCGT CAGCGAGGTGAGCAACGGCACGTCGAGGAGGGCAGGCTGCGTCTGGCTGCCTCTTCT GGTCTTGCACCTGCTTCTCAAATTTTGATGTGAGTGCCACTTCCCCACCCGGGAAAGGCT AAAGAATACTTTGGGGGGAAAAGAGTTTTAAAAAAAGAAATTGAAAATTGCCTTGCAGATA TTTAGGTACAATGGAGTTTTCTTTTCCCAAACGGGAAGAACACAGCACACCCGGCTTGGA CCCACTGCAAGCTGCATCGTGCAACCTCTTTGGTGCCAGTGTGGGCAAGGGCTCAGCCTC TCTGCCCACAGAGTGCCCCCACGTGGAACATTCTGGAGCTGGCCATCCCAAATTCAATCA GTCCATAGAGACGAACAGAATGAGACCTTCCGGCCCAAGCGTGGCGCTGCGGGCACTTTG

FIGURE 56

MKTIQPKMHNSISWAIFTGLAALCLFQGVPVRSGDATFPKAMDNVTVRQGESATLRCTID NRVTRVAWLNRSTILYAGNDKWCLDPRVVLLSNTQTQYSIEIQNVDVYDGGPYTCSVQTI NHPKTSRVHLIVQVSPKIVEISSDISINEGNNISLTCIATGREPFTVTWRHISPKAVGFV SEDEYLEIQGITREQSGDYECSASNDVAAPVVRRVKVTVNYPPYISEAKGTGVPVGQKGT LQCEASAVPSAEFQWYKDDKRLIEGKKGVKVENRPFLSKLIFFNVSEHDYGNYTCVASNK LGHTNASIMLFPGPAVSEVSNGTSRRAGCVWLDPLLVLHLLLKF

FIGURE 57

GCTGCGCCGGCTGCGGCTGCAGGGGAATCCGCTGTGCTGCGGCTGCCAGGCGCCCCCT CCTGCGGGCGAGGCTCTGGACGCCCTGCGGCCCTGGGACCTGCGCTGCCCTGGGGACGC GGCGCAGGAAGAGGAAGAGCTGGAAGAGCGGGCTGTGGCCGGGCCCCGCGCCCCTCCGCG $\tt CGGCCTCGCGCGGCCCCGGGGAGGAGCGGCAGTCGCCCTTGCCCTCGCGCCTGCGT$ GTGCGTCCCGAGTCCCGGCACAGCAGCTGCGAGGGCTGCGGCCTGCAGGCGGTGCCCCG CGGCTTCCCCAGCGACACCCAGCTCCTGGACCTGAGGCGGAACCACTTCCCCTCGGTGCC CCAGCGGCCTTCCCCGGNCTGGGCCACCTGGTGTCGCTGCACCTGCAGCACTGCGGCAT CGCGGAGCTGGAAGCGGGCGCCCTGGCCGGGCTGGGCCGCCTGATCTACCTGTACCTCTC CGACACCAGCTCGCAGGCCTCAGCGCTGCCCTTGAAGGGGCTCCCCGCCTCGGCTA CCTGTACCTAGAACGCAACCGTTTCCTGCAGGTGCCAGGGGCTGCCNTGCGCGCCCTGCC CAGCCTCTTCTCCCTGCACCTGCAGGACACGCTGTGGACCGCCTGGCACCTGGGGACCT GGGGAGAACACGGCCTTGCGCTGGGTCTACCTGAGTGGAAACCGCATCACCGAAGTGTC CCTTGGGGCGCTGGGCCCAGCTCGGGAGCTGGAGAAGCTGCACCTGGACAGGAATCAGCT GGGCAACCCACTCAGGGCCTTGCGTGACGGAGCCTTCCAGCCTGTGGGCAGGTCGCTGCA GCACCTCTTCCTGAACAGCAGTGGCCTGGAGCAGATTTGTCCTGGGGCCTTTTCAGGCCT GCCCAGTCTCAGCCAGCTGGAGCTCATCGACCTCAGCAGCAATCCCTTTCCCCTGTGACTG CCAGCTGCTTCCGCTGCACAGGTGGCTTACTGGGCTGAACCTGCGGGTGGGGGCCACCTG CGCCACCCTCCCAATGCCCGTGGCCAGAGGGTGAAGGCTGCAGCTGCTGTCTTTGAAGA CTGCCCGGGCTGGGCTGCCAGAAAGGCCAAGCGGACACCAGCCTCCAGGCCCAGTGCCAG GAGAACCCCCATCAAAGGAAGACAGTGTGGAGCAGATAAGAACATCCTCTTCCCCACATG GTACCACACTGTGGAGCCCACCTCGCTGTCATAGGCCTGCGGCTCTGAAGGATGGCTTTG CCCGCTCCCGCTCTGCCCCTCAAGTGGAACCCAAGCTGGGCTCAGAATCTGTAGAGTGAG GCCCACCAAGGGAAACGACACCCACGGCCTGAGAGCCAGGTGGAGTCCTGCCACTCAGC TGCCTGCCTTTGCTCCCACCCTCTCCCACCCTCAAAGAGGTCTCGAGGGGACACTCTGAA GGCACCTGGCTCAGAACCACTGCCATCCAAGGAGCGAGGAGTCCCAGGGCTGAGCAAATG CAGCGGGGAGGTCGCCAGTTCCCCTGCTTCCCGATCCTCATTTTCTGCTTCACTTGACTC CTCCAGATAGGAGCTGCTCTCACTGCCCACACTGCTG

FIGURE 58

LRRLRLQGNPLWCGCQARPLLEMLARARVRSDGACQGPRRLRGEALDALRPWDLRCPGDA
AQEEEELEERAVAGPRAPPRGPBRGPGEERAVAPCPRACVCVPESRHSSCEGCGLQAVPR
GFBSDTQLLDLRRNHFPSVPRAAFPGLGHLVSLHLQHCG1ABLEAGALAGLGRLIYLYLS
DNQLAGLSAAALEGAPRLGYLYLERNRFLQVPGAAXRALPSLFSLHLQDNAVDRLAPGDI
GRTRALRWYYLSGNRITEVSLGALGPARELEKHLDENDQLREVPTGALEGLPALLELQLS
GNPLRALRDGAFQPVGRSLQHLFLNSSGLEQICPGAFSGLGPGLQSLHLQKNQLRAPPAL
PSLSQLELIDLSSNPFPCDCQLLPLHRWLTGLNLRVGATCATPPNARGQRVKAAAAVFED
CPGWAARKAKTPASRPSARRTPIKGRQCGADKNILFPTWYHTVEPTSLS

Signal sequence:

None

Transmembrane domain:

None

N-glycosylation site:

325-328

Glycosaminoglycan attachment site: 338-341

Protein kinase C phosphorylation site: 438-440

N-myristoylation site:

166-171, 186-191, 253-258, 286-291, 335-340, 339-344, 450-455

Leucine rich repeat N-terminal domain: 94-123

Leucine Rich Repeat:

125-148, 149-172, 173-196, 197-220, 221-244, 245-268, 269-292, 293-316, 318-341, 343-364, 365-386

Leucine rich repeat C-terminal domain: 374-422

FIGURE 59

 $\tt CTCCCACGGTGTCCAGCGCCCAGA{\color{blue}ATGCGGCTTCTGGTCCTGCTATGGGGTTGCCTGCTG}$ CTCCCAGGTTATGAAGCCCTGGAGGGCCCAGAGGAAATCAGCGGGTTCGAAGGGGACACT GTGTCCTGCAGTGCACCTACAGGGAAGAGCTGAGGGACCACCGGAAGTACTGGTGCAGG AAGGGTGGGATCCTCTCTCTCGCTGCTCTGGCACCATCTATGCAGAAGAAGAAGGCCAG GAGACAATGAAGGGCAGGGTGTCCATCCGTGACAGCCGCCAGGAGCTCTCGCTCATTGTG ACCCTGTGGAACCTCACCCTGCAAGACGCTGGGGAGTACTGGTGTGGGGTCGAAAAACGG GGCCCGATGAGTCTTTACTGATCTCTCTGTTCGTCTTTCCAGGACCCTGCTGTCCTCCC TCCCCTTCTCCCACCTTCCAGCCTCTGGCTACAACACGCCTGCAGCCCAAGGCAAAAAGCT CAGCAAACCCAGCCCCAGGATTGACTTCTCCTGGGCTCTACCCGGCAGCCACCACAGCC A AGCAGGGGA AGACAGGGGCTGAGGCCCCTCCATTGCCAGGGACTTCCCAGTACGGCAC GAAAGGACTTCTCAGTACACAGGAACCTCTCCTCACCCAGCGACCTCTCCTCCTGCAGGG AGCTCCCCCCCCCATGCAGCTGGACTCCACCTCAGCAGAGGACACCAGTCCAGCTCTC AGCAGTGGCAGCTCTAAGCCCAGGGTGTCCATCCCGATGGTCCGCATACTGGCCCCAGTC CTGGTGCTGAGCCTTCTGTCAGCCGCAGGCCTGATCGCCTTCTGCAGCCACCTGCTC CTGTGGAGAAAGGAAGCTCAACAGGCCACGGAGACACAGAGGAACGAGAAGTTCTGGCTC TCACGCTTGACTGCGGAGGAAAAGGAAGCCCCTTCCCAGGCCCCTGAGGGGGACGTGATC TCGATGCCTCCCTCCACACATCTGAGGAGGAGCTGGGCTTCTCGAAGTTTGTCTCAGCG ACCGATTCCCGAAAGCTTTCCACCTCAGCCTCAGAGTCCAGCTGCCCGGACTCCAGGGCT CTCCCCACCTCCCCAGGCTCTCCTCTTGCATGTTCCAGCCTGACCTAGAAGCGTTTGTC AGCCTGGAGCCCAGAGCGGTGGCCTTGCTCTTCCGGCTGGAGACTGGGACATCCCTGAT AGGTTCACATCCCTGGGCAGAGTACCAGGCTGCTGACCCTCAGCAGGGCCAGACAAGGCT CAGTGGATCTGGTCTGAGTTTCAATCTGCCAGGAACTCCTGGGCCTCATGCCCAGTGTCG TTAGTCCCACGGTCTCCTGCATCAGCTGGTGATGAAGAGGAGCATGCTGGGGTGAGACTG GGATTCTGGCTTCTCTTTGAACCACCTGCATCCAGCCCTTCAGGAAGCCTGTGAAAAACG TGATTCCTGGCCCCACCAAGACCCACCAAAACCATCTCTGGGCTTGGTGCAGGACTCTGA ATTCTAACAATGCCCAGTGACTGTCGCACTTGAGTTTGAGGGCCAGTGGGCCTGATGAAC GCTCACACCCCTTCAGCTTAGAGTCTGCATTTGGGCTGTGACGTCTCCACCTGCCCCAAT AGATCTGCTCTGCCGACACCAGATCCACGTGGGGACTCCCCTGAGGCCTGCTAAGTC CAGGCCTTGGTCAGGTCAGGTGCACATTGCAGGATAAGCCCAGGACCGGCACAGAAGTGG TTGCCTTTNCCATTTGCCCTCCCTGGNCCATGCCTTCTTGCCTTTGGAAAAATGATGAA GAAAACCTTGGCTCCTTCCTTGTCTGGAAAGGGTTACTTGCCTATGGGTTCTGGTGGCTA GAGAGAAAAGTAGAAAACCAGAGTGCACGTAGGTGTCTAACACAGAGGAGAGTAGGAACA GGGCGGATACCTGAAGGTGACTCCGAGTCCAGCCCCCTGGAGAAGGGGTCGGGGGTGGTG GTAAAGTAGCACAACTACTATTTTTTTTTTTTTCCATTATTATTGTTTTTTAAGACAGA ATCTCGTGCTGCCCCAGGCTGGAGTGCAGTGGCACGATCTGCAAACTCCGCCTCCTGG ACACCTGGCTAATTTTTGTACTTTTAGTAGAGATGGGGTTTCACCATGTTGGCCAGGCTG GTCTTGAACTCCTGACCTCAAATGAGCCTCCTGCTTCAGTCTCCCAAATTGCCGGGATTA CAGGCATGAGCCACTGTGTCTGGCCCTATTTCCTTTAAAAAGTGAAATTAAGAGTTGTTC AGTATGCAAAACTTGGAAAGATGGAGGAGAAAAAGAAAAGGAAGAAAAAAATGTCACCCA TTCACATAATTTGCCGGTGTTCTTTTTACAGAGCAATTATCTTGTATATACAACTTTGTA TCCTGCCTTTTCCACCTTATCGTTCCATCACTTTATTCCAGCACTTCTCTGTGTTTTACA

FIGURE 60

MRLLVLLWGCLLLPGYEALEGPEEISGFEGDTVSLQCTYREELRDHRKYWCRKGGILFSR CSGTIYAEEEGQETMKGRVSIRDSRGELSLIVTLWNLTLQDAGEYWGGVEKRGPDESLLI SLFVPFBCPCCPPSFSPTFOPLATTRLQPKAKAQQTOPFGLTSFGLYPAATTAKQGKTGAE APPLPGTSQYGHERTSQYTGTSPHPATSPPAGSSRPPMQLDSTSAEDTSPALSSGSSKPR VSIPMVRILAPVLVLLSLLSAAGLIAFCSHLLLWRKEAQQATETQRNEKFWLSRLTAEEK EAPSOAPEGDVISMPPLHTSEEELGFSKFVSA

Important features: Signal peptide: amino acids 1-17

Transmembrane domain: amino acids 248-269

N-glycosylation site: amino acids 96-99

Fibrinogen beta and gamma chains C-terminal domain: amino acids 104-113

Ig like V-type domain: amino acids 13-128

FIGURE 61

CGGGCCAGCCTGGGGCGGCCGGCCAGGAACCACCCGTTAAGGTGTCTTCTCTTTAGGGAT GGTGAGGTTGGAAAAAGACTCCTGTAACCCTCCTCCAGGATGAACCACCTGCCAGAAGAC ATGGAGAACGCTCTCACCGGGAGCCAGAGCTCCCATGCTTCTCTGCGCAATATCCATTCC ATATCTGATGTCAGGAGGACTTTCTGTTTGTTTGTCACCTTTGACCTCTTATTCGTAACA TTACTGTGGATAATAGAGTTAAATGTGAATGGAGGCATTGAGAACACATTAGAGAAGGAG GTGATGCAGTATGACTACTATTCTTCATATTTTGATATATTTCTTCTGGCAGTTTTTCGA TTTAAAGTGTTAATACTTGCATATGCTGTGTGCAGACTGCGCCATTGGTGGGCAATAGCG TTGACAACGGCAGTGACCAGTGCCTTTTTACTAGCAAAAGTGATCCTTTCGAAGCTTTTC ACGTGGTTCCTGGATTTCAAAGTGTTACCTCAAGAAGCAGAAGAAGAAAACAGACTCCTG ATAGTTCAGGATGCTTCAGAGAGGGCAGCACTTATACCTGGTGGTCTTTCTGATGGTCAG TTTTATTCCCCTCTGAATCCGAAGCAGGATCTGAAGAAGCTGAAGAAAAACAGGACAGT GAGAAACCACTTTTAGAACTATGAGTACTACTTTTGTTAAATGTGAAAAACCCTCACAGA AAGTCATCGAGGCAAAAAGAGGCAGGCAGTGGAGTCTCCCTGTCGACAGTAAAGTTGAAA TGTCATGATTCATCCTCTCTCAGTGAGACTGAGCCTGATGTGTTAACAAATAGGTGAAG AAAGTCTTGTGCTGTATTCCTAATCAAAAGACTTAATATATTGAAGTAACACTTTTTTAG ATTTATTTGTATTTCTTTTTTAACACTCTACATTTCCCTTGTTTTTTAACTCATGCACA TGTGCTCTTTGTACAGTTTTAAAAAGTGTAATAAAATCTGACATGTCAATGTGGCTAGTT TTATTTTTCTTGTTTTGCATTATGTGTATGGCCTGAAGTGTTGGACTTGCAAAAGGGGAA GAAAGGAATTGCGAATACATGTAAAATGTCACCAGACATTTGTATTATTTTTATCATGAA ATCATGTTTTTCTCTGATTGTTCTGAAATGTTCTAAATACTCTTATTTTGAATGCACAAA ATGACTTAAACCATTCATATCATGTTTCCTTTGCGTTCAGCCAATTTCAATTAAAATGAA CTAAATTAAAAA

FIGURE 62

MNHLPEDMENALTGSQSSHASLRNIHSINPTQLMARIESYEGREKKGISDVRRTFCLFVT FDLLFVTLLWIIELNVNGGIENTLEKEVMQYDYYSSYFDIFLLAVFRFKVLILAYAVCRL RHWWAIALTTAVTSAFLLAKVILSKLFSQGAFGYVLPIISFILAWIETWFLDFKVLPQEA EEENRLLIVODASERAALIPGGLSDGQFYSPPESEAGSEEAEEKQDSEKPLLEL

Important features of the protein: Signal peptide: amino acids 1-20

Transmembrane domains: amino acids 54-72, 100-118, 130-144, 146-166

N-myristoylation sites: amino acids 14-20, 78-84, 79-85, 202-208, 217-223

FIGURE 63

ATGGCTGCAGCCACCTCGCGCGCACCCCGAGGCGCCCCGCGCCCAGCTCGCCCGAGGTCCGT CGGAGGCGCCGGCCCCCGGAGCCAAGCAGCAACTGAGCGGGGAAGCGCCCGCGTCCG GGGATCGGGATGTCCCTCCTCCTCCTCTCTCTGCTAGTTTCCTACTATGTTGGAACCTTG GGGACTCACACTGAGATCAAGAGAGTGGCAGAGGAAAAGGTCACTTTGCCCTGCCACCAT CAACTGGGGCTTCCAGAAAAAGACACTCTGGATATTGAATGGCTGCTCACCGATAATGAA GGGAACCAAAAAGTGGTGATCACTTACTCCAGTCGTCATGTCTACAATAACTTGACTGAG GAACAGAAGGGCCGAGTGGCCTTTGCTTCCAATTTCCTGGCAGGAGATGCCTCCTTGCAG ATTGAACCTCTGAAGCCCAGTGATGAGGGCCGGTACACCTGTAAGGTTAAGAATTCAGGG CGCTACGTGTGGAGCCATGTCATCTTAAAAGTCTTAGTGAGACCATCCAAGCCCAAGTGT GAGTTGGAAGGAGGCTGACAGAAGGAAGTGACCTGACTTTGCAGTGTGAGTCATCCTCT GGCACAGAGCCCATTGTGTATTACTGGCAGCGAATCCGAGAGAAAGAGGGAGAGGATGAA CGTCTGCCTCCCAAATCTAGGATTGACTACAACCACCCTGGACGAGTTCTGCTGCAGAAT AGCTGTGTGGTGCGAGTAACTGTACAGTATGTACAAAGCATCGGCATGGTTGCAGGAGCA GTGACAGGCATAGTGGCTGGAGCCCTGCTGATTTTCCTCTTGGTGTGGCTGCTAATCCGA AGGAAAGACAAAGAAAGATATGAGGAAGAAGAGAGACCTAATGAAATTCGAGAAGATGCT GAAGCTCCAAAAGCCCGTCTTGTGAAACCCAGCTCCTCTTCCTCAGGCTCTCGGAGCTCA CTGTCAACTGACGCAGCACCCCAGCCAGGCTGGCCACCCAGGCATACAGCCTAGTGGGG CCAGAGGTGAGAGGTTCTGAACCAAAGAAAGTCCACCATGCTAATCTGACCAAAGCAGAA ACCACACCCAGCATGATCCCCAGCCAGAGCAGAGCCTTCCAAACGGTC**TGA**ATTACAATG GACTTGACTCCCACGCTTTCCTAGGAGTCAGGGTCTTTGGACTCTTCTCGTCATTGGAGC TCAAGTCACCAGCCACACAACCAGATGAGAGGTCATCTAAGTAGCAGTGAGCATTGCACG GAACAGATTCAGATGAGCATTTTCCTTATACAATACCAAACAAGCAAAAGGATGTAAGCT GATTCATCTGTAAAAAGGCATCTTATTGTGCCTTTAGACCAGAGTAAGGGAAAGCAGGAG TCCAAATCTATTTGTTGACCAGGACCTGTGGTGAGAAGGTTGGGGAAAGGTGAGGTGAAT ATACCTAAAACTTTTAATGTGGGATATTTTGTATCAGTGCTTTGATTCACAATTTTCAAG AGGAAATGGGATGCTGTTTGTAAATTTTCTATGCATTTCTGCAAACTTATTGGATTATTA GTTATTCAGACAGTCAAGCAGAACCCACAGCCTTATTACACCTGTCTACACCATGTACTG AGCTAACCACTTCTAAGAAACTCCAAAAAAGGAAACATGTGTCTTCTATTCTGACTTAAC TTCATTTGTCATAAGGTTTGGATATTAATTTCAAGGGGAGTTGAAATAGTGGGAGATGGA GAAGAGTGAATGAGTTTCTCCCACTCTATACTAATCTCACTATTTGTATTGAGCCCAAAA TAACTATGAAAGGAGACAAAAATTTGTGACAAAGGATTGTGAAGAGCTTTCCATCTTCAT GATGTTATGAGGATTGTTGACAAACATTAGAAATATATAATGGAGCAATTGTGGATTTCC CCTCAAATCAGATGCCTCTAAGGACTTTCCTGCTAGATATTTCTGGAAGGAGAAAATACA ACATGTCATTTATCAACGTCCTTAGAAAGAATTCTTCTAGAGAAAAAGGGATCTAGGAAT AATTGCAAGACTGGGTGGACTAGAAAGGGAGATTAGATCAGTTTTCTCTTAATATGTCAA GGAAGGTAGCCGGGCATGGTGCCAGGCACCTGTAGGAAAATCCAGCAGGTGGAGGTTGCA GTGAGCCGAGATTATGCCATTGCACTCCAGCCTGGGTGACAGAGCGGGACTCCGTCTC

FIGURE 64

MSLLLLLLVSYYVGTLGTHTEIKRVAEEKVTLPCHHQLGLPEKDTLDIEWLLTDNEGNQ
KVVITYSSRHVYNNLTEEQKGRVAFASNFLAGDASLQIEPLKPSDEGRYTCKVKNSGRYV
WSHVILKVLVRPSKPKCELEGELTEGSDLTLQCESSSGTEPIVYYWQRIREKEGEDERLP
PKSRIDYNHPGRVLLQNLTMSYSGLYQCTAGNEAGKESCVVRVTVQYVQSIGMVAGAVTG
IVAGALLIFLLVWLLIRRKDKERYEEEERPNEIREDAEAPKARLVKPSSSSGSRSSRSG
SSSTRSTANASARSQRTLSTDAAPQPGLATQAYSLVGPEVRGSEPKKVHHANLTKAETTP
SMIPSOSRAFOTV

Signal sequence:

Transmembrane domain: amino acids 232-251

FIGURE 65

GTCGGGGCTGCGCGACGGCGCAGGGGCTGCGGGGAGCGCCGCGCAGGCCGTGCAGTTCCT AGCGAGGAGGCGCCGCCATTGCCGCTCTCTCGGTGAGCGCAGCCCCGCTCTCCGGGC CGGGCCTTCGCGGGCCACCGGCGCCATGGGCCAGTGCGGCATCACCTCCTCCAAGACCGT GCTGGTCTTTCTCAACCTCATCTTCTGGGGGGGCAGCTGGCATTTTATGCTATGTGGGAGC CTATGTCTTCATCACTTATGATGACTATGACCACTTCTTTGAAGATGTGTACACGCTCAT CCCTGCTGTAGTGATCATAGCTGTAGGAGCCCTGCTTTTCATCATTGGGCTAATTGGCTG CTGTGCCACAATCCGGGAAAGTCGCTGTGGACTTGCCACGTTTGTCATCATCCTGCTCTT GGTTTTTGTCACAGAAGTTGTTGTAGTGGTTTTGGGATATGTTTACAGAGCAAAGGTGGA TGCTGCTAGCCGGGCTATTGATTATGTACAGAGACAGCTGCATTGTTGTGGAATTCACAA CTACTCAGACTGGGAAAATACAGATTGGTTCAAAGAAACCAAAAACCAGAGTGTCCCTCT TAGCTGCTGCAGAGAGACTGCCAGCAATTGTAATGGCAGCCTGGCCCACCCTTCCGACCT CTATGCTGAGGGGTGTGAGGCTCTAGTAGTGAAGAAGCTACAAGAAATCATGATGCATGT GATCTGGGCCGCACTGGCATTTGCAGCTATTCAGCTGCTGGGCATGCTGTGCTTGCAT CGTGTTGTGCAGAAGGAGTAGAGATCCTGCTTACGAGCTCCTCATCACTGGCGGAACCTA TGCATAGTTGACAACTCAAGCCTGAGCTTTTTGGTCTTGTTCTGATTTGGAAGGTGAATT GAGCAGGTCTGCTGCTGTTGGCCTCTGGAGTTCATTTAGTTAAAGCACATGTACACTGGT GTTGGACAGAGCAGCTTGGCTTTTCATGTGCCCACCTACTTACCTACTACCTGCGACTTT CTTTTTCCTTGTTCTAGCTGACTCTTCATGCCCCTAAGATTTTAAGTACGATGGTGAACG TTCTAATTTCAGAACCAATTGCGAGTCATGTAGTGTGGTAGAATTAAAGGAGGACACGAG CCTGCTTCTGTTACCTCCAAGTGGTAACAGGACTGATGCCGAAATGTCACCAGGTCCTTT CAGTCTTCACAGTGGAGAACTCTTGGCCAAAGGTTTTTGCGGGGAGGAGGAGGAAACCAG CTTTCTGGTTAAGGTTAACACCAGATGGTGCCCCTCATTGGTGTCCTTTTAAAAAATATT TACTGTAGTCCAATAAGATAGCAGCTGTACAAAATGACTAAAATAGATTGTAGGATCATA TGGCGTATATCTTGGTTCATCTTCAAAATCAGAGACTGAGCTTTGAAACTAGTGGTTTTT AATCAAAGTTGGCTTTATAGGAGGAGTATAATGTATGCACTACTGTTTTAAAAGAATTAG TGTGAGTGTGTTTTTGTATGAATGAGCCCATTCATGGTAAGTCTTAAGCTTGTTGGAAAT AATGTACCCATGTAGACTAGCAAAATAGTATGTAGATGTGATCTCAGTTGTAAATAGAAA

FIGURE 66

MGQCGITSSKTVLVFLNLIFWGAAGILCYVGAYVFITYDDYDHFFEDVYTLIPAVVIIAV GALLFIIGLIGCCATIRESRCGLATFVIILLLVFVTEVVVVVLGYVYRAKVENEVDRSIQ KVYKTYNGTNPDAASRAIDYVQRQLHCCGIHNYSDWENTDWFKETKNQSVPLSCCRETAS NCNGSLAHPSDLYAEGCEALVVKKLQEIMMHVIWAALAFAAIQLLGMLCACIVLCRRSRD PAYFILITGGTYA

Signal peptide:

Type II transmembrane domain: 11-38

Other transmembrane domains: 48-68, 87-107, 208-235

N-glycosylation site: 127-131, 152-156, 167-171, 183-187

Tyrosine kinase phosphorylation site: 236-244

N-myristoylation site: 5-11, 68-74, 71-77, 226-232

Prokaryotic membrane lipoprotein lipid attachment site: 62-73, 221-232

Transmembrane 4 family proteins: 7-35, 56-106

FIGURE 67

 $\tt GCGGCACCTGGAAG\underline{A}TGCGCCCATTGGCTGGTGGCCTGCTCAAGGTGGTGTTCGTGGTCT$ TCGCCTCCTTGTGTGCCTGGTATTCGGGGTACCTGCTCGCAGAGCTCATTCCAGATGCAC CCCTGTCCAGTGCTGCCTATAGCATCCGCAGCATCGGGGAGAGGCCTGTCCTCAAAGCTC CAGTCCCCAAAAGGCAAAAATGTGACCACTGGACTCCCTGCCCATCTGACACCTATGCCT ACAGGTTACTCAGCGGAGGTGGCAGAAGCAAGTACGCCAAAATCTGCTTTGAGGATAACC TACTTATGGGAGAACAGCTGGGAAATGTTGCCAGAGGAATAAACATTGCCATTGTCAACT GACCGATGACAAAGTTTATTCAGAGTGCTGCTCCAAAATCCCTGCTCTTCATGGTGACCT ATGACGACGGAAGCACAAGACTGAATAACGATGCCAAGAATGCCATAGAAGCACTTGGAA TGGAACTCCCTTCCGAAATTCAGAGAGAAAAGATCAACCACTCTGATGCTAAGAACAACA GATATTCTGGCTGGCCTGCAGAGATCCAGATAGAAGGCTGCATACCCAAAGAACGAAGCT GACACTGCAGGGTCCTGAGTAAATGTGTTCTGTATAAACAAATGCAGCTGGAATCGCTCA AGAATCTTATTTTCTAAATCCAACAGCCCATATTTGATGAGTATTTTGGGTTTGTTGTA AACCAATGAACATTTGCTAGTTGTATCAAATCTTGGTACGCAGTATTTTTATACCAGTAT AAA

FIGURE 68

MRPLAGGLLKVVFVVFASLCAWYSGYLLAELIPDAPLSSAAYSIRSIGERPVLKAPVPKR QKCDHWTPCPSDTYAYRLLSGGGRSKYAKICFEDNLLMGEQLGNVARGINIAIVNYVTGN VTATRCFDMYEGDNSGPMTKFIQSAAPKSLLFMVTYDDGSTRLNNDAKNAIEALGSKEIR NMKFRSSWVFIAAKGLELPSEIQREKINHSDAKNNRYSGWPAEIQIEGCIPKERS

Signal sequence: amino acids 1-20

N-glycosylation sites: amino acids 120-124, 208-212

Glycosaminoglycan attachment site: amino acids 80-84

N-myristoylation sites: amino acids 81-87, 108-114, 119-125

FIGURE 69

ACACA ACTTTA CACCTGAATGAACGCCAAACCTCTATGGATATATAAAGGGAAGCTTGAG GAGGAATTTCACAGTTACAGTGCAGAAGCAGAAGCAAAAGAATTAACCAGCTCTTCAGTC CATGCATCCTAATGAAAAGCTGTTTGGCTTTTAAAAAATGATGCCACAGAAATCCTTTATT CACATGTGGTTAAACCTGTTCCAGCACACCCCAGCAGCAACAGCACGTTGAATCAAGCCA GAAATGGAGGCAGGCATTTCAGTAACACTGGACTGGATCGGAACACTCGGGTTCAAGTGG GTTGCCGGGAACTGCGTTCCACCAAATACATCTCTGATGGCCAGTGCACCAGCATCAGCC CTCTGAAGGAGCTGGTGTGTGCTGGCGAGTGCTTGCCCCTGCCAGTGCTCCCTAACTGGA TTGGAGGAGGCTATGGAACAAAGTACTGGAGCAGGAGGAGCTCCCAGGAGTGGCGGTGTG TCAATGACAAAACCCGTACCCAGAGAATCCAGCTGCAGTGCCAAGATGGCAGCACACGCA CCTACAAAATCACAGTAGTCACTGCCTGCAAGTGCAAGAGGTACACCCGGCAGCACAACG AGTCCAGTCACAACTTTGAGAGCATGTCACCTGCCAAGCCAGTCCAGCATCACAGAGAGC GGAAAAGAGCCAGCAAATCCAGCAAGCACAGCATGAGT<u>TAG</u>AACTCAGACTCCCATAACT AGACTTACTAGTAACCATCTGCTTTACAGATTTGATTGCTTGGAAGACTCAAGCCTGCCA CTGCTGTTTTCTCACTTGAAAGTATATGCTTTCTGCTTTGATCAAACCCAGCAAGCTGTC TTAAGTATCAGGACCTTCTTTGGGAATAGTTTTTCCTTTTAAAGTTTTTCAAGATGTAGG TATATCCATGAATGCAATTTGCATTTAAATTCCACGTATCCCTGTAGTTTAAATTCCTCA TTGGTCTTAAAAGACTGTTGATACTATAAACATCAGTGGAATCAATTATATTTTAAAACA GAAAAGGGCTT

FIGURE 70

MLPPAIHFYLLPLACILMKSCLAFKNDATEILYSHVVKPVPAHPSSNSTLNQARNGGRHF SNTGLDRNTRVQVGCRELRSTKYISDGQCTSISPLKELVCAGECLPLPVLPNWIGGGYGT KYWSRRSSQEWRCVNDKTRTQRIQLQCQDGSTRTYKITVVTACKCKRYTRQHNESSHNFE SMSPAKPVOHHRERKRASKSSKHSMS

Signal sequence: 1-23

Transmembrane domain:

N-glycosylation site: 47-50, 173-176

cAMP- and cGMP-dependent protein kinase phosphorylation site: 125-128, 166-169, 195-198

N-myristoylation site: 64-69, 87-92, 115-120, 116-121, 150-155

FIGURE 71

CCCAGGCTCTAGTGCAGGAGGAGAAGGAGGAGGAGGAGGAGGTGGAGATTCCCAGTTAAA AGGCTCCAGAATCGTGTACCAGGCAGAGAACTGAAGTACTGGGGCCTCCTCCACTGGGTC CGAATCAGTAGGTGACCCCGCCCCTGGATTCTGGAAGACCTCACCATGGGACGCCCCCGA CCTCGTGCGGCCAAGACGTGGATGTTCCTGCTCTTGCTGGGGGGGAGCCTGGGCAGGACAC TCCAGGGCACAGGAGACAAGGTGCTGGGGGGTCATGAGTGCCAACCCCATTCGCAGCCT TGGCAGGCGCCTTGTTCCAGGGCCAGCAACTACTCTGTGGCGGTGTCCTTGTAGGTGGC AACTGGGTCCTTACAGCTGCCCACTGTAAAAAACCGAAATACACAGTACGCCTGGGAGAC CACAGCCTACAGAATAAAGATGGCCCAGAGCAAGAAATACCTGTGGTTCAGTCCATCCCA CACCCTGCTACAACAGCAGCGATGTGGAGGACCACAACCATGATCTGATGCTTCTTCAA CTGCGTGACCAGGCATCCCTGGGGTCCAAAGTGAAGCCCATCAGCCTGGCAGATCATTGC ACCCAGCCTGGCCAGAAGTGCACCGTCTCAGGCTGGGGCACTGTCACCAGTCCCCGAGAG AATTTTCCTGACACTCTCAACTGTGCAGAAGTAAAAATCTTTCCCCAGAAGAAGTGTGAG GACACGTGCCAGGGCGATTCTGGAGGCCCCCTGGTGTGTGATGGTGCACTCCAGGGCATC ACATCCTGGGGCTCAGACCCCTGTGGGAGGTCCGACAAACCTGGCGTCTATACCAACATC ${\tt TGCCGCTACCTGGACTGGATCAAGAAGATCATAGGCAGCAAGGGC\underline{{\tt TGA}}{\tt TTCTAGGATAAG}$ CACTAGATCTCCCTTAATAAACTCACAACTCTCTGGTTC

FIGURE 72

MGRPRPRAAKTWMFLLLLGGAWAGHSRAQEDKVLGGHECQPHSQPWQAALFQGQQLLCGG VLVGGNWVLTAAHCKKPKYTVRLGDHSLQNKDGPEQEIPVVQSIPHPCYNSSDVEDHNHD LMLLQLRDQASLGSKVKPISLADHCTQPGQKCTVSGWGTVTSPRENFPDTLNCAEVKIFP QKKCEDAYPGQITDGMVCAGSSKGADTCQGDSGGPLVCDGALQGITSWGSDPCGRSDKPG VYTNICRYLDWIKKIIGSKG

Important Features: Signal peptide: amino acids 1-23

Transmembrane domain: amino acids 51-71

N-glycosylation site: amino acids 110-113

Serine proteases, trypsin family, histidine active site: amino acids 69-74 and 207-217

Tyrosine kinase phosphorylation site: amino acids 182-188

Kringle domain proteins motif: amino acids 205-217

FIGURE 73

CCGGCCTCTCCAATGGCAAATGTGTGTGGCTGGAGGCGAGGCGAGGCTTTCGGCAAAGG CAGTCGAGTGTTTGCAGACCGGGGCGAGTCCTGTGAAAGCAGATAAAAGAAAACATTTAT TAACGTGTCATTACGAGGGGGGCCCCGGCCGGGGCTGTCGCACTCCCCGCGGAACATTT GGCTCCCTCCAGCTCCGAGAGAGAGAAGAAGAAAGCGGAAAAAGAGGCAGATTCACGTCG TTTCCAGCCAAGTGGACCTGATCGATGGCCCTCCTGAATTTATCACGATATTTGATTTAT CTTCCCTCCCTCGTTTCCAGCTCCTGGGCGAATCCCACATCTGTTTCAACTCTCCGCCGA GGGCGAGCAGGAGCGAGAGTGTGTCGAATCTGCGAGTGAAGAGGGACGAGGGAAAAGAAA CAAAGCCACAGACGCAACTTGAGACTCCCGCATCCCAAAAGAAGCACCAGATCAGCAAAA AAAGAAGATGGGCCCCCCGAGCCTCGTGCTGTGCTTGCTGTCCGCAACTGTGTTCTCCCT GCTGGGTGGAAGCTCGGCCTTCCTGTCGCACCACCGCCTGAAAGGCAGGTTTCAGAGGGA GGGTTCCATGCAGGTGATGAACAAGACCCGGCGCATCATGGAGCAGGGCGGGGCGCACTT CATCAACGCCTTCGTGACCACACCCATGTGCTGCCCCTCACGCTCCTCCATCCTCACTGG CAAGTACGTCCACAACCACAACACCTACACCAACAATGAGAACTGCTCCTCGCCCTCCTG GCAGGCACAGCACGAGAGCCGCACCTTTGCCGTGTACCTCAATAGCACTGGCTACCGGAC AGCTTTCTTCGGGAAGTATCTTAATGAATACAACGGCTCCTACGTGCCACCCGGCTGGAA GGAGTGGGTCGGACTCCTTAAAAACTCCCGCTTTTATAACTACACGCTGTGTCGGAACGG GGTGAAAGAGAAGCACGGCTCCGACTACTCCAAGGATTACCTCACAGACCTCATCACCAA TGACAGCGTGAGCTTCTTCCGCACGTCCAAGAAGATGTACCCGCACAGGCCAGTCCTCAT GGTCATCAGCCATGCAGCCCCCACGGCCCTGAGGATTCAGCCCCACAATATTCACGCCT CTTCCCAAACGCATCTCAGCACATCACGCCGAGCTACAACTACGCGCCCAACCCGGACAA ACACTGGATCATGCGCTACACGGGGCCCATGAAGCCCATCCACATGGAATTCACCAACAT GCTCCAGCGGAAGCGCTTGCAGACCCTCATGTCGGTGGACGACTCCATGGAGACGATTTA CAACATGCTGGTTGAGACGGGCGAGCTGGACAACACGTACATCGTATACACCGCCGACCA CGGTTACCACATCGGCCAGTTTGGCCTGGTGAAAGGGAAATCCATGCCATATGAGTTTGA CATCAGGGTCCCGTTCTACGTGAGGGGCCCCAACGTGGAAGCCGGCTGTCTGAATCCCCA CATCGTCCTCAACATTGACCTGGCCCCCACCATCCTGGACATTGCAGGCCTGGACATACC TGCGGATATGGACGGGAAATCCATCCTCAAGCTGCTGGACACGGAGCGGCCGGTGAATCG GTTTCACTTGAAAAAGAAGATGAGGGTCTGGCGGGACTCCTTCTTGGTGGAGAGAGGCAA GCTGCTACACAAGAGAGACAATGACAAGGTGGACGCCCAGGAGGAGAACTTTCTGCCCAA GTACCAGCGTGTGAAGGACCTGTGTCAGCGTGCTGAGTACCAGACGGCGTGTGAGCAGCT GGGACAGAAGTGGCAGTGTGTGGAGGACGCCACGGGGAAGCTGAAGCTGCATAAGTGCAA GGGCCCCATGCGGCTGGGCGGCAGCAGAGCCCTCTCCAACCTCGTGCCCAAGTACTACGG GCAGGGCAGCGAGGCCTGCACCTGTGACAGCGGGGACTACAAGCTCAGCCTGGCCGGACG CCGGAAAAAACTCTTCAAGAAGAAGTACAAGGCCAGCTATGTCCGCAGTCGCTCCATCCG CTCAGTGGCCATCGAGGTGGACGGCAGGGTGTACCACGTAGGCCTGGGTGATGCCGCCCA GCCCCGAAACCTCACCAAGCGGCACTGGCCAGGGGCCCCTGAGGACCAAGATGACAAGGA TGGTGGGGACTTCAGTGGCACTGGAGGCCTTCCCGACTACTCAGCCGCCAACCCCATTAA AGTGACACATCGGTGCTACATCCTAGAGAACGACACAGTCCAGTGTGACCTGGACCTGTA CAAGTCCCTGCAGGCCTGGAAAGACCACAAGCTGCACATCGACCACGAGATTGAAACCCT GCAGAACAAAATTAAGAACCTGAGGGAAGTCCGAGGTCACCTGAAGAAAAAGCGGCCAGA AGAATGTGACTGTCACAAAATCAGCTACCACACCCAGCACAAAGGCCGCCTCAAGCACAG AGGCTCCAGTCTGCATCCTTTCAGGAAGGGCCTGCAAGAGAAGGACAAGGTGTGGCTGTT GCGGGAGCAGAAGCGCAAGAAAACTCCGCAAGCTGCTCAAGCGCCTGCAGAACAACGA CACGTGCAGCATGCCAGGCCTCACGTGCTTCACCCACGACAACCAGCACTGGCAGACGGC

GCCTTTCTGGACACTGGGGCCTTTCTGTGCCTGCACCAGCGCCAACAATAACACGTACTG GTGCATGAGGACCATCAATGAGACTCACAATTTCCTCTTCTGTGAATTTGCAACTGGCTT CCTAGAGTACTTTGATCTCAACACAGACCCCTACCAGCTGATGAATGCAGTGAACACACT GGACAGGGATGTCCTCAACCAGCTACACGTACAGCTCATGGAGCTGAGGAGCTGCAAGGG ATACAGGCAGTTTCAGCGTCGAAAGTGGCCAGAAATGAAGAGACCTTCTTCCAAATCACT TAGAGGCATCACCTGACTGCACAGGCAATGAAAAACCATGTGGGTGATTTCCAGCAGACC TCTGGAGGATAACCAGCAGGAGCAGAGATAACTTCAGGAAGTCCATTTTTGCCCCTGCTT TTGCTTTGGATTATACCTCACCAGCTGCACAAAATGCATTTTTTCGTATCAAAAAGTCAC CTTGGAAATTTCTCCCAAGGGCGAAAGTCATTGGAATTTTTAAATCATAGGGGAAAAGCA ACAGAGGCAACGTGGAGAGGCTGAAAACAGTGCAGAGACGTTTGACAATGAGTCAGTAGC ACAAAAGAGATGACATTTACCTAGCACTATAAACCCTGGTTGCCTCTGAAGAAACTGCCT TCATTGTATATATGTGACTATTTACATGTAATCAACATGGGAACTTTTAGGGGAACCTAA TAAGAAATCCCAATTTTCAGGAGTGGTGGTCAATAAACGCTCTGTGGCCAGTGTAAAA GAAAAA

FIGURE 74

MGPPSLVLCLLSATVFSLLGGSSAFLSHHRLKGRFQRDRRNIRPNIILVLTDDQDVELGS
MQVMNKTRRIMEQGGAHFINAFVTTPMCCPSRSSILTGKYVHNHNTYTMENCSSPSWQA
QHESRTFAVYLNSTGYRTAFFGKYLNEYNGSTVPFGMKEWVGLLKNSRFYNYTLCRNGVK
EKHGSDYSKDYLTDLITNDSVSFFRTSKKMYPHRPVLMVISHAAPHGPEDSAPQYSRLFP
NASQHITPSYNYAPNPDKHWIMRYTGFMKPIHMEFTIMMLOKRRLQTLMSVDDSMETIYNM
LVETGELDNTYIVYTADHGYHIGOFGLVKGKSMPYEFDIRVPFYVRGPNVEAGCLMPHIV
LNIDLAPTILDIAGLDIPADMDGKSILKLLDTERPVMRFHLKKKMRWMRDSFLVERGKLL
KKRDNDKVDAQEENFLPKYQRVKDLCQRAEYQTACEQLGKWQCVEDATGKLKLHKCKGP
MRLGGSRALSNLVPKYYGGGSEACTCDSGDYKLSLAGRRKKLFKKKYKASYVRSRSIRSV
AIEVDGRVYHVGLGDAAQPRNLTKRHWPGAPEDQDDKDGGDFSGTGGLPDYSAANPIKVT
HRCYILENDTVQCDLDLYKSLQAWKDHKLHIDHEIETLQNKIKNLREVRGHLKKKRPEC
DCHKISYHTQHKGRIKHRGSSLHPFRKGLQEKDKVWLLREQKRKKLKKKLKLLKRLQNNDTC
SMPGLTCFTHDNQHWQTAPFWTLGPFCACTSANNNTYWCMRTINETHNFLFCEFATGFLE
YFDLNTDFVQLMNAVNTLDRDVLNQLHVQLMELRSCKGYKQCNPRTRNMDLDGGSYEQYR

Important features: Signal peptide: amino acids 1-17

Sulfatases signature 1: amino acids 86-99

Homologous region to sulfatase: amino acids 87-106, 133-146, 216-229, 291-320, 365-375

N-glycosylation sites: amino acids 65-69, 112-116, 132-136, 149-153, 171-175, 198-202, 241-245, 561-565, 608-612, 717-721, 754-758, 764-768

FIGURE 75

CCCACGCGTCCGCCCACGCGTCCGGTGGACTATGGGCCAGTTTTTGTGCAAGAACCAGAT GATATTATTTTTCCAACTGATTCTGATGAAAAGAAGGTAGCATTGAATTGTGAAGTTCGT GGCAATCCAGTTCCCAGTTACAGATGGCTTCGAAATGGAACAGAAATAGATCTGGAAAGT GATTATCGCTACAGTTTGATAGATGGCACCTTCATTATAAGCAATCCAAGTGAAGCAAAG GATTCTGGTCATTATCAGTGTTTAGCAACCAACACTGTGGGGAGTATTCTTAGTAGAGAA GCTACACTGCAGTTTGCCTATCTGGGAAATTTTAGTGGCCGGACAAGAAGTGCAGTCTCT GTGAGGGAAGGCCAGGGTGTCGTTCTGATGTGCTCTCCTCCGCCACATTCACCAGAGATC ATCTATAGCTGGGTATTTAATGAGTTCCCTTCCTTTGTGGCGGAAGACAGCCGGCGGTTC ATCTCCCAGGAGACAGGCAACCTTTATATTTCTAAAGTCCAAACATCAGATGTTGGCAGC TATATTTGTCTGGTGAAAAACACAGTGACGAATGCTAGAGTCCTTAGTCCTCCAACGCCA $\tt CTCACTCTGCGTAATGATGGTGTGATGGGAGAATATGAGCCGAAAATTGAGGTCCATTTT$ AACCCCGTTCCAACAATCACATGGATGAAGGTTAATGGTTATATTCCTAGTAAGGCACGT CTGCGGAAATCTCAGGCGGTGCTGGAAATACCGAATGTACAGCTGGATGATGCAGGCATT TATGAGTGCAGAGCTGAAAACTCACGTGGAAAAAATTCCTTTCGTGGACAATTACAAGTA TACACCTACCCACACTGGGTAGAAAAACTGAATGATACTCAGTTAGACAGTGGGAGCCCT CTCCGATGGGAATGTAAGGCTACTGGAAAACCCAGACCCACGTATCGTTGGCTGAAGAAT GGAGTACCCCTCTCACCTCAGAGTAGGGTTGAGATGGTTAATGGAGTATTGATGATCCAC AATGTGAATCAATCAGATGCTGGAATGTATCAGTGTTTGGCTGAAAATAAGTATGGAGCC ATTTACGCTAGTGCTGAGCTGAAGATTCTAGCTTCAGCTCCCACTTTTGCACTGAATCAA CTGAAGAAACAATAATTGTTACCAAAGACCAAGAAGTTGTCATAGAGTGCAAACCCCAA GGCTCTCCAAAACCAACCATCTCTTGGAAGAAAGGAGACAGAGCAGTTAGAGAAAACAAA AGAATAGCTATTCTTCCAGACGGGAGTCTACGGATCCTAAATGCTTCCAAATCAGACGAG GGAAAGTACGTTTGCCGAGGGGAAAACGTCTTTGGTTCTGCTGAAAT

FIGURE 76

MCSPPPHSPEIIYSWVFNEFPSFVAEDSRRFISQETGNLYISKVQTSDVGSYICLVKNTV
TNARVLSPPTPLTLRNDGVMGEYEPKLEVHFFFTVTAAKGTTVKMECFALGNPVPTITWM
KVNGYIPSKARLRKSQAVLEIPNVQLDDAGIYECRAENSRGKNSFRGQLQVYTYPHWVEK
LNDTQLDSGSPLRWECKATGKPRPTYRWLKNGVPLSPQSRVEMVNGVLMIHNVNQSDAGM
YQCLAENKYGAIYASAELKILASAPTFALNQLKKTITVTKDQEVVIECKPQGSPKPTISW
KKGDRAVRENKRIAILPDGSLRTLINASKSDEGKYVCRGENVFGSAE

Signal sequence:

None

Transmembrane domain:

None

N-glycosylation site: 182-185, 234-237, 325-328

Tyrosine kinase phosphorylation site: 328-334

N-myristoylation site: 50-55, 150-155, 239-244, 250-255

Immunoglobulin domain: 2-56, 100-156, 189-245, 281-338

FIGURE 77

GCTCCCAGCCAAGAACCTCGGGGCCGCTGCGCGGTGGGGAGGAGTTCCCCGAAACCCGGC CGCTAAGCGAGGCCTCCTCCTCCCGCAGATCCGAACGGCCTGGGCGGGGTCACCCCGGCT GGGACAAGAAGCCGCCGCCTGCCTGCCCGGGCCCGGGGAGGGGGCTGGGGCTGGGGCCGG AGGCGGGGTGTGAGTGGGTGTGTGCGGGGGGGGGGGGCTTGATGCAATCCCGATAAGAAA TGCTCGGGTGTCTTGGGCACCTACCCGTGGGGCCCGTAAGGCGCTACTATATAAGGCTGC CGGCCCGGAGCCGCCGCCGCCGTCAGAGCAGGAGCGCTGCGTCCAGGATCTAGGGCCACGA CCATCCCAACCCGGCACTCACAGCCCCGCAGCGCATCCCGGTCGCCGCCCAGCCTCCCGC ACCCCATCGCCGGAGCTGCGCCGAGAGCCCCAGGGAGGTGCC<u>AT</u>GCGGAGCGGGTGTGT GGTGGTCCACGTATGGATCCTGGCCGGCCTCTGGCTGGCCGTGGCCGGGCGCCCCCTCGC CTTCTCGGACGCGGGCCCCACGTGCACTACGGCTGGGGCGACCCCATCCGCCTGCGGCA CCTGTACACCTCCGGCCCCCACGGGCTCTCCAGCTGCTTCCTGCGCATCCGTGCCGACGG CGTCGTGGACTGCGCGCGGGGCCAGAGCGCGCACAGTTTGCTGGAGATCAAGGCAGTCGC TCTGCGGACCGTGGCCATCAAGGGCGTGCACAGCGTGCGGTACCTCTGCATGGGCGCCGA CGGCAAGATGCAGGGGCTGCTTCAGTACTCGGAGGAAGACTGTGCTTTCGAGGAGGAGAT CCGCCCAGATGGCTACAATGTGTACCGATCCGAGAAGCACCGCCTCCCGGTCTCCCTGAG CAGTGCCAAACAGCGGCAGCTGTACAAGAACAGAGGCTTTCTTCCACTCTCATTTCCT GCCCATGCTGCCCATGGTCCCAGAGGAGCCTGAGGACCTCAGGGGCCACTTGGAATCTGA CATGTTCTCTCGCCCCTGGAGACCGACAGCATGGACCCATTTGGGCTTGTCACCGGACT GGAGGCCGTGAGGAGTCCCAGCTTTGAGAAGTAACTGAGACCATGCCCGGGCCTCTTCAC TGCTGCCAGGGGCTGTGGTACCTGCAGCGTGGGGGACGTGCTTCTACAAGAACAGTCCTG AGTCCACGTTCTGTTTAGCTTTAGGAAGAAACATCTAGAAGTTGTACATATTCAGAGTTT TCCATTGGCAGTGCCAGTTTCTAGCCAATAGACTTGTCTGATCATAACATTGTAAGCCTG TAGCTTGCCCAGCTGCTGGCCCGGGCCCCCATTCTGCTCCCTCGAGGTTGCTGGACAAGCT GCTGCACTGTCTCAGTTCTGCTTGAATACCTCCATCGATGGGGAACTCACTTCCTTTGGA AAAATTCTTATGTCAAGCTGAAATTCTCTAATTTTTTCTCATCACTTCCCCAGGAGCAGC CAGAAGACAGGCAGTAGTTTTAATTTCAGGAACAGGTGATCCACTCTGTAAAACAGCAGG TAAATTTCACTCAACCCCATGTGGGAATTGATCTATATCTCTACTTCCAGGGACCATTTG CCCTTCCCAAATCCCTCCAGGCCAGAACTGACTGGAGCAGGCATGGCCCACCAGGCTTCA GGAGTAGGGGAAGCCTGGAGCCCCACTCCAGCCCTGGGACAACTTGAGAATTCCCCCTGA GGCCAGTTCTGTCATGGATGCTGTCCTGAGAATAACTTGCTGTCCCGGTGTCACCTGCTT CCATCTCCCAGCCCACCAGCCCTCTGCCCACCTCACATGCCTCCCCATGGATTGGGGCCT CCCAGGCCCCCACCTTATGTCAACCTGCACTTCTTGTTCAAAAATCAGGAAAAGAAAAG ATTTGAAGACCCCAAGTCTTGTCAATAACTTGCTGTGTGGAAGCAGCGGGGGAAGACCTA GAGGTTTGTTTTGTATATTAAAATGGAGTTTGTTTGT

FIGURE 78

MRSGCVVVHVWILAGLWLAVAGRPLAFSDAGPHVHYGWGDPIRLRHLYTSGPHGLSSCFL RIRADGVVDCARGQSAHSLLEIKAVALRTVAIKGVHSVRYLCMGADGKMQGLLQYSEEDC AFEEEIRPDGYNVYRSEKHRLPVSLSSAKQRQLYKNRGFLPLSHFLPMLPMVPEEPEDLR GHLESDMFSSPLETDSMDFFGLVTGLEAVRSPSFEK

Signal peptide: amino acids 1-22

Casein kinase II phosphorylation site: amino acids 78-82, 116-120, 190-194, 204-208

N-myristoylation site: amino acids 15-21, 54-60, 66-72, 201-207

Prokaryotic membrane lipoprotein lipid attachment site: amino acids 48-59

FIGURE 79

CGGACGCGTGGCCGACGCGTGGGCCTGGGCAAGGGCCGGGGCCGGGCCGAGCCACCTCTTCCC CTCCCCCGCTTCCCTGTCGCGCTCCGCTGGCTGGACGCGCTGGAGGAGTGGAGCACCCCGGCCG GCCCTGGGGGCTGACAGTCGGCAAAGTTTGGCCCGAAGAGGGAAGTGGTCTCAAACCCCGGCAGGTG TGGCATCCAGAGTACGGGTCGAGCCCGGGCCATGGAGCCCCCCTGGGGAGGCGGCACCAGGGAGCC TGGGCGCCGGGGCTCCGCCGCGACCCCATCGGGTAGACCACAGAAGCTCCGGGACCCTTCCGGCA CCTCTGGACAGCCCAGGATGCTGTTGGCCACCCTCCTCCTCCTCCTCCTTGGAGGCGCTCTGGCCC ATCCAGACCGGATTATTTTTCCAAATCATGCTTGTGAGGACCCCCCAGCAGTGCTCTTAGAAGTGC AGGGCACCTTACAGAGGCCCCTGGTCCGGGACAGCCGCACCTCCCCTGCCAACTGCACCTGGCTCA TCCTGGGCAGCAAGGAACAGACTGTCACCATCAGGTTCCAGAAGCTACACCTGGCCTGTGGCTCAG AGCGCTTAACCCTACGCTCCCCTCTCCAGCCACTGATCTCCCTGTGTGAGGCACCTCCCAGCCCTC TGCAGCTGCCCGGGGGCAACGTCACCATCACTTACAGCTATGCTGGGGCCAGAGCACCCATGGGCC AGGGCTTCCTGCTCCTACAGCCAAGATTGGCTGATGTGCCTGCAGGAAGAGTTTCAGTGCCTGA ACCACCGCTGTGTATCTGCTGTCCAGCGCTGTGATGGGGTTGATGCCTGTGGCGATGGCTCTGATG ATGTCACCTTGGAGGACTTCTATGGGGTCTTCTCCTCTCTGGATATACACACCTAGCCTCAGTCT CCCACCCCAGTCCTGCCATTGGCTGCTGGACCCCCATGATGGCCGGCGGCTGGCCGTGCGCTTCA CAGCCCTGGACTTGGAGATGCAGTGCATGTGTATGACGGCCCTGGGCCCCTGAGAGCT CCCGACTACTGCGTAGTCTCACCCACTTCAGCAATGGCAAGGCTGTCACTGTGGAGACACTGTCTG GCCAGGCTGTTGTGTCCTACCACACAGTTGCTTGGAGCAATGGTCGTGGCTTCAATGCCACCTACC ATGTGCGGGGCTATTGCTTGCCTTGGGACAGACCCTGTGGCTTAGGCTCTGGCCTGGGAGCTGGCG AAGGCCTAGGTGAGCGCTGCTACAGTGAGGCACAGCGCTGTGACGGCTCATGGGACTGTGCTGACG GCACAGATGAGGAGGACTGCCCAGGCTGCCCACCTGGACACTTCCCCTGTGGGGCTGCTGGCACCT CTGGTGCCACAGCCTGCTACCTGCCTGCTGACCGCTGCAACTACCAGACTTTCTGTGCTGATGGAG CAGATGAGAGACGCTGTCGGCATTGCCAGCCTGGCAATTTCCGATGCCGGGACGAGAAGTGCGTGT ATGAGACGTGGGTGTGCGATGGGCAGCCAGACTGTGCGGACGGCAGTGATGAGTGGGACTGCTCCT ATGTTCTGCCCCGCAAGGTCATTACAGCTGCAGTCATTGGCAGCCTAGTGTGCGGCCTGCTCCTGG TCATCGCCCTGGGCTGCACCTGCAAGCTCTATGCCATTCGCACCCAGGAGTACAGCATCTTTGCCC CCCTCTCCCGGATGGAGGCTGAGATTGTGCAGCAGCAGCACCCCCTTCCTACGGGCAGCTCATTG CCCAGGGTGCCATCCCACCTGTAGAAGACTTTCCTACAGAGAATCCTAATGATAACTCAGTGCTGG GCAACCTGCGTTCTCTGCTACAGATCTTACGCCAGGATATGACTCCAGGAGGTGGCCCAGGTGCCC GCCGTCGTCAGCGGGCCGCTTGATGCGACGCCTGGTACGCCGTCTCCGCCGCTGGGGCTTGCTCC CTCGAACCAACACCCCGGCTCGGGCCTCTGAGGCCAGATCCCAGGTCACACCTTCTGCTGCTCCCC AGCAGGCACCCCCACTGCCCATCAAGGCTCCCCTCCCATCTGCTAGCACGTCTCCAGCCCCCACTA CTGTCCCTGAAGCCCCAGGGCCACTGCCCTCACTGCCCCTAGAGCCATCACTATTGTCTGGAGTGG TGCAGGCCCTGCGAGGCCGCCTGTTGCCCAGCCTGGGGCCCCCAGGACCAACCCGGAGCCCCCCTG GACCCCACACAGCAGTCCTGGCCCTGGAAGATGAGGACGATGTGCTACTGGTGCCACTGGCTGAGC TCCACCACTTCCTTCCCTGTCCCTGGATTTCAGGGACTTGGTGGGCCTCCCGTTGACCCTATGTAG CTGCTATAAAGTTAAGTGTCCCTCAGGCAGGGAGAGGGCTCACAGAGTCTCCTCTGTACGTGGCCA TGGCCAGACACCCCAGTCCCTTCACCACCACCTGCTCCCCACGCCACCACCATTTGGGTGGCTGTT TTTAAAAAGTAAAGTTCTTAGAGGATCATAGGTCTGGACACTCCATCCTTGCCAAACCTCTACCCA AAAGTGGCCTTAAGCACCGGAATGCCAATTAACTAGAGACCCTCCAGCCCCCAAGGGGAGGATTTG GGCAGAACCTGAGGTTTTGCCATCCACAATCCCTCCTACAGGGCCTGGCTCACAAAAAAGAGTGCAA CAAATGCTTCTATTCCATAGCTACGGCATTGCTCAGTAAGTTGAGGTCAAAAATAAAGGAATCATA CATCTC

FIGURE 80

MLLATLLLLLGGALAHPDRIIFPNHACEDPPAVLLEVQGTLQRPLVRDSRTSPANCTWL
ILGSKEQTTTIRFÇKLHLACGSERLTLRSPLQPLISLCEAPPSPLQLPGGNVTITYSYAG
ARAPMGQGFLLSYSQDWLMCLQEEFQCLNHRCVSAVQRCDGVDACGGSDEBGCGSDPFP
GLTPRPVPSLPCNVTLEDFYGVFSSPGYTHLASVSHPQSCHULDPHDGRRLAVRFTALD
LGFGDAVHVYDGPGPPESSRLLRSLTHFSNGKAVTVETLSGQAVVSYHTVAWSNGRGFNA
TYHVRGYCLPWDRPGCLGSGLGAGGGGGGECYSEAGRCDGSWDCADGTDEEDCPGCPPGH
FPCGAAGTSGATACYLPADRCNYQTFCADGADERRCRHCQPGNFRCRDEKCYYETWVCDG
QPDCADGSDEWDCSYVLPRKVITAAVIGSLVCGLLVIALGCTCKLYAIRTQEYSIFAPL
SRMEAEIVQQQAPPSYGGLIAQGAIPPVEDFPTENPNDNSVLGNLRSLLQILRQDMTPGG
GPGARRRQRGRLMRRLVRRLRRWGLLPRTNTPARASEARSQVTPSAAPLEALDGGTGPAR
EGGAVGGQDGEQAPPLPIKAPLSSATSPAPTTVPEAPGPLPSLPLEPSLLSGVVQALRG
RLLPSLGPPGPTRSPPGPHTAVLALEDEDDVLLVPLAEPGWWVAEAEDEPLLT

Important features:
Signal peptide:
amino acids 1-16

Transmembrane domain:

LDL-receptor class A (LDLRA) domain proteins: amino acids 411-431, 152-171, 331-350 and 374-393

FIGURE 81

CTTCTGTGCTGTTCCTTCTTGCCTCTAACTTGTAAACAAGACGTACTAGGACGATGCTAA TGGAAAGTCACAAACCGCTGGGTTTTTGAAAGGATCCTTGGGACCTCATGCACATTTGTG GAAACTGGATGGAGAGTTTGGGGAAGCATGGACTCTTTAGCCAGCTTAGTTCTCTGTGG AGTCAGCTTGCTCCTTTCTGGAACTGTGGAAGGTGCCATGGACTTGATCTTGATCAATTC CCTACCTCTTGTATCTGATGCTGAAACATCTCTCACCTGCATTGCCTCTGGGTGGCGCCC CCATGAGCCCATCACCATAGGAAGGGACTTTGAAGCCTTAATGAACCAGCACCAGGATCC GCTGGAAGTTACTCAAGATGTGACCAGAGAATGGGCTAAAAAAGTTGTTTGGAAGAGAGA AAAGGCTAGTAAGATCAATGGTGCTTATTTCTGTGAAGGGCGAGTTCGAGGAGAGCCAAT GACTGTGGACAAGGGAGATAACGTGAACATATCTTTCAAAAAGGTATTGATTAAAGAAGA AGATGCAGTGATTTACAAAAATGGTTCCTTCATCCATTCAGTGCCCCGGCATGAAGTACC TGATATTCTAGAAGTACACCTGCCTCATGCTCAGCCCCAGGATGCTGGAGTGTACTCGGC CAGGTATATAGGAGGAAACCTCTTCACCTCGGCCTTCACCAGGCTGATAGTCCGGAGATG TGAAGCCCAGAAGTGGGGACCTGAATGCAACCATCTCTGTACTGCTTGTATGAACAATGG TGTCTGCCATGAAGATACTGGAGAATGCATTTGCCCTCCTGGGTTTATGGGAAGGACGTG AGAGGGATGCAAGTCTTATGTGTTCTGTCTCCCTGACCCCTATGGGTGTTCCTGTGCCAC AGGCTGGAAGGGTCTGCAGTGCAATGAAGCATGCCACCCTGGTTTTTACGGGCCAGATTG ${\tt TAAGCTTAGGTGCAGCTGCAACAATGGGGAGATGTTGTGATCGCTTCCAAGGATGTCTCTG}$ CTCTCCAGGATGGCAGGGGCTCCAGTGTGAGAGAGAGGCATACCGAGGATGACCCCAAA GATAGTGGATTTGCCAGATCATATAGAAGTAAACAGTGGTAAATTTAATCCCATTTGCAA AGCTTCTGGCTGGCCGCTACCTACTAATGAAGAAATGACCCTGGTGAAGCCGGATGGGAC AGTGCTCCATCCAAAAGACTTTAACCATACGGATCATTTCTCAGTAGCCATATTCACCAT CCACCGGATCCTCCCCCTGACTCAGGAGTTTGGGTCTGCAGTGTGAACACAGTGGCTGG GATGGTGGAAAAGCCCTTCAACATTTCTGTTAAAGTTCTTCCAAAGCCCCTGAATGCCCC AAACGTGATTGACACTGGACATAACTTTGCTGTCATCAACATCAGCTCTGAGCCTTACTT TGGGGATGGACCAATCAAATCCAAGAAGCTTCTATACAAACCCGTTAATCACTATGAGGC TTGGCAACATATTCAAGTGACAAATGAGATTGTTACACTCAACTATTTGGAACCTCGGAC AGAATATGAACTCTGTGTGCAACTGGTCCGTCGTGGAGAGGGTGGGGAAGGGCATCCTGG CCTGCCTAAAAGTCAGACCACTCTAAATTTGACCTGGCAACCAATATTTCCAAGCTCGGA AGATGACTTTTATGTTGAAGTGGAGAGAAGGTCTGTGCAAAAAAGTGATCAGCAGAATAT TAAAGTTCCAGGCAACTTGACTTCGGTGCTACTTAACAACTTACATCCCAGGGAGCAGTA CGTGGTCCGAGCTAGAGTCAACACCAAGGCCCAGGGGGAATGGAGTGAAGATCTCACTGC TTGGACCCTTAGTGACATTCTTCCTCCTCAACCAGAAAACATCAAGATTTCCAACATTAC ACACTCCTCGGCTGTGATTTCTTGGACAATATTGGATGGCTATTCTATTTCTTCTATTAC TATCCGTTACAAGGTTCAAGGCAAGAATGAAGACCAGCACGTTGATGTGAAGATAAAGAA TGCCACCATCATTCAGTATCAGCTCAAGGGCCTAGAGCCTGAAACAGCATACCAGGTGGA CATTTTTGCAGAGAACAACATAGGGTCAAGCAACCCAGCCTTTTCTCATGAACTGGTGAC CCTCCCAGAATCTCAAGCACCAGCGGACCTCGGAGGGGGGAAGATGCTGCTTATAGCCAT CCTTGGCTCTGCTGGAATGACCTGCCTGACTGTGCTGTTGGCCTTTCTGATCATATTGCA ATTGAAGAGGGCAAATGTGCAAAGGAGAATGGCCCAAGCCTTCCAAAACGTGAGGGAAGA ACCAGCTGTGCAGTTCAACTCAGGGACTCTGGCCCTAAACAGGAAGGTCAAAAACAACCC AGATCCTACAATTTATCCAGTGCTTGACTGGAATGACATCAAATTTCAAGATGTGATTGG GGAGGGCAATTTTGGCCAAGTTCTTAAGGCGCGCATCAAGAAGGATGGGTTACGGATGGA TGCTGCCATCAAAAGAATGAAAGAATATGCCTCCAAAGATGATCACAGGGACTTTGCAGG AGAACTGGAAGTTCTTTGTAAACTTGGACACCATCCAAACATCATCAATCTCTTAGGAGC

ATGTGAACATCGAGGCTACTTGTACCTGGCCATTGAGTACGCGCCCCATGGAAACCTTCT GGACTTCCTTCGCAAGAGCCGTGTGCTGGAGACGGACCCAGCATTTGCCATTGCCAATAG CACCGCGTCCACACTGTCCTCCCAGCAGCTCCTTCACTTCGCTGCCGACGTGGCCCGGGG CATGGACTACTTGAGCCAAAAACAGTTTATCCACAGGGATCTGGCTGCCAGAAACATTTT AGTTGGTGAAAACTATGTGGCAAAAATAGCAGATTTTGGATTGTCCCGAGGTCAAGAGGT GTACGTGAAAAAGACAATGGGAAGGCTCCCAGTGCGCTGGATGGCCATCGAGTCACTGAA TTACAGTGTGTACACAACCAACAGTGATGTATGGTCCTATGGTGTTTACTATGGGAGAT TGTTAGCTTAGGAGGCACACCCTACTGCGGGATGACTTGTGCAGAACTCTACGAGAAGCT GCCCCAGGGCTACAGACTGGAGAAGCCCCTGAACTGTGATGATGAGGTGTATGATCTAAT GAGACAATGCTGGCGGGAGAAGCCTTATGAGAGGCCATCATTTGCCCAGATATTGGTGTC CTTAAACAGAATGTTAGAGGAGCGAAAGACCTACGTGAATACCACGCTTTATGAGAAGTT ${\tt TACTTATGCAGGAATTGACTGTTCTGCTGAAGAAGCGGCC\underline{{\tt TAG}}{\tt GACAGAACATCTGTATA}$ CCCTCTGTTTCCCTTTCACTGGCATGGGAGACCCTTGACAACTGCTGAGAAAACATGCCT CTGCCAAAGGATGTGATATATAAGTGTACATATGTGCTGGAATTCTAACAAGTCATAGGT TCCCTCACCTGTAGCATGCCAGTCCCGTTTCATTTAGTCATGTGACCACTCTGTCTTGTG TTTCCACAGCCTGCAAGTTCAGTCCAGGATGCTAACATCTAAAAATAGACTTAAATCTCA TTGCTTACAAGCCTAAGAATCTTTAGAGAAGTATACATAAGTTTAGGATAAAATAATGGG ATTTTCTTTTCTCTGGTAATATTGACTTGTATATTTAAGAAATAACAGAAAGCC TGGGTGACATTTGGGAGACATGTGACATTTATATATTGAATTAATATCCCTACATGTATT GCACATTGTAAAAAGTTTTAGTTTTGATGAGTTGTGAGTTTACCTTGTATACTGTAGGCA

FIGURE 82

MDSLASLVLCGVSLLLSGTVEGAMDLILINSLPLVSDAETSLTCIASGWRPHEPITIGRD FEALMNOHODPLEVTODVTREWAKKVVWKREKASKINGAYFCEGRVRGEAIRIRTMKMRO OASFLPATLTMTVDKGDNVNISFKKVLIKEEDAVIYKNGSFIHSVPRHEVPDILEVHLPH AQPQDAGVYSARYIGGNLFTSAFTRLIVRRCEAQKWGPECNHLCTACMNNGVCHEDTGEC ICPPGFMGRTCEKACELHTFGRTCKERCSGOEGCKSYVFCLPDPYGCSCATGWKGLOCNE ACHPGFYGPDCKLRCSCNNGEMCDRFQGCLCSPGWQGLQCEREGIPRMTPKIVDLPDHIE VNSGKFNPICKASGWPLPTNEEMTLVKPDGTVLHPKDFNHTDHFSVAIFTIHRILPPDSG VWVCSVNTVAGMVEKPFNISVKVLPKPLNAPNVIDTGHNFAVINISSEPYFGDGPIKSKK LLYKPVNHYEAWOHIOVTNEIVTLNYLEPRTEYELCVOLVRRGEGGEGHPGPVRRFTTAS IGLPPPRGLNLLPKSOTTLNLTWOPIFPSSEDDFYVEVERRSVQKSDQQNIKVPGNLTSV LLNNLHPREOYVVRARVNTKAQGEWSEDLTAWTLSDILPPOPENIKISNITHSSAVISWT ILDGYSISSITIRYKVQGKNEDQHVDVKIKNATIIQYQLKGLEPETAYQVDIFAENNIGS SNPAFSHELVTLPESQAPADLGGGKMLLIAILGSAGMTCLTVLLAFLIILOLKRANVORR MAOAFONVREEPAVOFNSGTLALNRKVKNNPDPTIYPVLDWNDIKFQDVIGEGNFGQVLK ARIKKDGLRMDAAIKRMKEYASKDDHRDFAGELEVLCKLGHHPNIINLLGACEHRGYLYL AIEYAPHGNLLDFLRKSRVLETDPAFAIANSTASTLSSQQLLHFAADVARGMDYLSQKQF IHRDLAARNILVGENYVAKIADFGLSRGQEVYVKKTMGRLPVRWMAIESLNYSVYTTNSD VWSYGVLLWEIVSLGGTPYCGMTCAELYEKLPQGYRLEKPLNCDDEVYDLMRQCWREKPY ERPSFAQILVSLNRMLEERKTYVNTTLYEKFTYAGIDCSAEEAA

Signal sequence:

1-38

Transmembrane domain:

750-770

N-glycosylation site:

140-143, 158-161, 399-402, 438-441, 464-467, 560-563, 596-599, 649-652, 691-694, 930-933, 1011-1014, 1104-1107

cAMP- and cGMP-dependent protein kinase phosphorylation site: 534-537

Tyrosine kinase phosphorylation site: 149-156. 808-816. 1094-1102

N-myristoylation site: 18-23, 98-103, 187-192, 196-201, 270-275, 286-291, 295-300, 420-425, 595-600, 984-989, 1036-1041, 1041-1046, 1115-1120

Prokaryotic membrane lipoprotein lipid attachment site: 882-892

EGF-like domain cysteine pattern signature: 240-251, 287-298, 329-340

Tyrosine protein kinases specific active-site signature: 960-972

Protein kinase domain: 824-1092

Fibronectin type III domain: 444-529, 543-626, 639-724

EGF-like domain: 220-251, 268-298

laminin_EGF Laminin EGF-like (Domains III and V):
219-268

Immunoglobulin domain: 156-193

Zinc finger:

Receptor tyrosine kinase: 844-868, 869-898, 936-982, 986-1024, 1025-1052, 1052-1088

FIGURE 83

GAAAGCAGCGAGTTGGCAGAGCAGGGCTGCATTTCCAGCAGGAGCTGCGAGCACAGTGCT GGCTCACAACAAG<u>AT</u>GCTCAAGGTGTCAGCCGTACTGTGTGTGTGCAGCCGCTTGGTG TAATTTTCTGGATGATAAACAATGGCTCACCACAATCTCTCAGTATGACAAGGAAGTCGG ACAGTGGAACAAATTCCGAGACGAAGTAGAGGATGATTATTTCCGCACTTGGAGTCCAGG AAAACCCTTCGATCAGGCTTTAGATCCAGCTAAGGATCCATGCTTAAAGATGAAATGTAG TCGCCATAAAGTATGCATTGCTCAAGATTCTCAGACTGCAGTCTGCATTAGTCACCGGAG GCTTACACACAGGATGAAAGAAGCAGGAGTAGACCATAGGCAGTGGAGGGGTCCCATATT ATCCACCTGCAAGCAGTGCCCAGTGGTCTATCCCAGCCCTGTTTGTGGTTCAGATGGTCA TACCTACTCTTTTCAGTGCAAACTAGAATATCAGGCATGTGTCTTAGGAAAACAGATCTC AGTCAAATGTGAAGGACATTGCCCATGTCCTTCAGATAAGCCCACCAGTACAAGCAGAAA TGTTAAGAGAGCATGCAGTGACCTGGAGTTCAGGGAAGTGGCAAACAGATTGCGGGACTG GTTCAAGGCCCTTCATGAAAGTGGAAGTCAAAACAAGAAGACAAAAACATTGCTGAGGCC TGAGAGAAGCAGATTCGATACCAGCATCTTGCCAATTTGCAAGGACTCACTTGGCTGGAT GTTTAACAGACTTGATACAAACTATGACCTGCTATTGGACCAGTCAGAGCTCAGAAGCAT TTACCTTGATAAGAATGAACAGTGTACCAAGGCATTCTTCAATTCTTGTGACACATACAA GGACAGTTTAATATCTAATAATGAGTGGTGCTACTGCTTCCAGAGACAGCAAGACCCACC TTGCCAGACTGAGCTCAGCAATATTCAGAAGCGGCAAGGGGTAAAGAAGCTCCTAGGACA GTATATCCCCCTGTGTGATGAAGATGGTTACTACAAGCCAACACAATGTCATGGCAGTGT TGGACAGTGCTGGTGTTGACAGATATGGAAATGAAGTCATGGGATCCAGAATAAATGG TGTTGCAGATTGTGCTATAGATTTTGAGATCTCCGGAGATTTTGCTAGTGGCGATTTTCA ${f T}{f CA}{f T}{f T}{f G}{f A}{f T}{f G}{f A}{f C}{f A}{f A}{f A}{f A}{f T}{f C}{f A}{f A}{f A}{f A}{f T}{f C}{f A}{f C}{f A}{f C}{f A}{f A}{f$ CCTATTTAAAATTATCTTCTCCCCAATAACAAAATGATTCTAAACCTCACATATATTTT GTATAATTATTTGAAAAATTGCAGCTAAAGTTATAGAACTTTATGTTTAAATAAGAATCA TTTGCTTTGAGTTTTTATATTCCTTACACAAAAAGAAAATACATATGCAGTCTAGTCAGA CAAAATAAAGTTTTGAAGTGCTACTATAATAAATTTTTCACGAGAACAAACTTTGTAAAT CTTCCATAAGCAAAATGACAGCTAGTGCTTGGGATCGTACATGTTAATTTTTTGAAAGAT AATTCTAAGTGAAATTTAAAATAAATAAATTTTTAATGACCTGGGTCTTAAGGATTTAGG AAAAATATGCATGCTTTAATTGCATTTCCAAAGTAGCATCTTGCTAGACCTAGATGAGTC

FIGURE 84

MLKVSAVLCVCAAAWCSQSLAAAAAVAAAGGRSDGGNFLDDKQWLTTISQYDKEVGQWNK
FRDEVEDDYFRTWSPGKPFDQALDPAKDPCLKMKCSRHKVCIAQDSQTAVCISHRRLTHR
MKEAGVDHRQWRGPILSTCKQCPUVYPSPVCGSDGHTYSFQCKLEYQACVLGKQISVKCE
GHCPCPSDKPTSTSRNVKRACSDLEFREVANRLRDWFKALHESGSQNKKTKTLLRPERSF
FDTSILPICKDSLGWMFNRLDTNYDLLLDQSELRSIYLDKNBGCTKAFFNSCDTYKDSLI
SNNEWCYCFQRQQDPPCQTELSNIQKRQGVKKLLGQYIPLCDEDGYYKPTQCHGSVGQCW
CVDRYGNEVMGSRINGVADCAIDFEISGDFASGDFHEWTDDEDDEDDIMNDEDEIEDDDE
DGGDDDDGGDDHDYVI

Important features: Signal peptide: amino acids 1-16

Leucine zipper pattern: amino acids 246-267

N-myristoylation sites: amino acids 357-362, 371-376 and 376-381

Thyroglobulin type-1 repeat proteins: amino acids 353-365 and 339-352

FIGURE 85

CCCACGCGTCCGGCACTGCAGTCTCCAGCCTGAGCCATGGGCCGCCGAGCCCTCCTGCTC TCGGTTCTCTACTTCCAACAGAAGGTTGATCATTTTGGATTTAATACTGTGAAAACTTTT AATCAGCGGTACCTAGTAGCTGATAAATACTGGAAGAAAAATGGTGGATCAATACTTTTC TACACTGGTAATGAAGGGGACATTATCTGGTTTTGTAATAACACGGGGTTCATGTGGGAT GTGGCTGAGGAACTGAAAGCTATGTTGGTGTTTGCTGAACATCGATACTATGGAGAGTCT CTCCCCTTTGGTGACAACTCATTCAAGGATTCCAGACACTTGAATTTCCTGACATCAGAA CAAGCTCTGGCTGATTTTGCAGAGTTAATCAAACACTTGAAAAGAACAATCCCAGGAGCT GAAAATCAACCTGTCATTGCCATAGGAGGCTCCTATGGTGGCATGCTTGCCGCCTGGTTT AGGATGAAATATCCTCATATGGTAGTTGGAGCTCTTGCAGCTTCTGCCCCTATCTGGCAG TTTGAGGATTTAGTACCTTGTGGTGTATTTATGAAGATCGTAACTACAGATTTTAGGAAA AGCGGTCCACATTGTTCAGAGAGCATCCACAGGTCCTGGGATGCCATTAATCGACTCTCA AATACTGGCAGTGGTTTGCAGTGGCTTACTGGAGCCCTTCACTTATGCAGCCCATTAACT TCTCAGGACATCCAACATTTGAAAGACTGGATCTCTGAAACCTGGGTGAATCTGGCAATG GTGGACTATCCTTATGCCTCTAACTTTTTACAGCCTTTGCCTGCTTGGCCTATCAAGGTA GTGTGCCAGTATTTGAAAAATCCCAATGTATCTGATTCACTGCTGCTGCAGAATATTTTC CAAGCTCTGAATGTATATTACAATTATTCGGGCCAGGTGAAATGCCTGAATATTTCAGAG ACAGCAACTAGCAGTCTGGGAACACTGGGTTGGAGCTATCAGGCCTGCACAGAAGTAGTC GAACTTTCTGATGACTGTTTTCAACAGTGGGGTGTGAGACCAAGGCCCTCCTGGATCACT ACTATGTATGGAGGCAAAAACATTAGTTCACACACAAACATTGTTTTCAGCAATGGTGAA CTAGACCCCTGGTCAGGAGGTGGAGTAACTAAGGATATCACAGACACTCTGGTTGCAGTC ACCATCTCAGAGGGGGCCCACCACTTAGATCTCCGCACCAAGAATGCCTTGGATCCTATG TCTGTGCTGTTAGCCCGCTCCTTGGAAGTTAGACATATGAAGAATTGGATCAGAGATTTC TATGACAGTGCGGGAAAGCAGCACTGAGAAACTTTTGATTGTTTTCAATTTCTTCTTTTA TGTTCACACCACCACATTCCCATTCACTTTGATTTTCTACATGTAATTACCTTCTTTTGT TTATCATTAGATTTGATGGGGCCAAAGTTGAGATAGAATAGAGGGTGATGACGGTAAGAG CAAGTGTCCCATGAATGTGATTTCCTGGGTTCTCACTGTCCTTTGCACCACGTCTAGGAA GAATCTTCTTGATAGCTCTCCCACACCATCAGTGGCCCTCATAACTGGAGTAGAGTTCCT GGTTGCTTTTCATAAGAGGGAGAGTTACTTTC

FIGURE 86

MGRRALLLLLSFLAPWATIALRPALRALGSLHLPTNPTSLPAVAKNYSVLYFQQKVDHF
GFNTVKTFNQRYLVADKYWKKNGGS LIFYTGNEGDIIWFGNNTGFMWDVAEELKAMLVFA
EHRYYGESLPFGDNSFKDSRHLNFLTSEQALADFAELIKHLKRTI PGAENQPVIAIGGSY
GGMLAAWFRMKYPHMVVGALAASAPIWQFEDLVPCGVFMKIVTTDFRKSGPHCSESIHRS
WDAINRLSNTGSGLQWLTGALHLCSPLTSQDIQHLKDWISETWVNLAMVDYPYASNFLQP
LPAMPIKVVCQYLKNPNVSDSLLLQNIFQALMVYYNYSGQVKCLNISETATSSLGTLGWS
YQACTEVVMPFCTNGVDDMFEPHSWNLKELSDDCFQQWGVRPRPSWITTMYGGKNISSHT
NIVPSNGELDPWGGGGVTKDITDTLVAVTISEGAHHLDLRTKNALDPMSVLLARSLEVRH
MKNWIRDFYDSAGKQH

```
Signal sequence:
1-18

Transmembrane domain:
None

N-glycosylation site:
47-50, 101-104, 317-320, 336-339, 345-348, 415-418
Glycosaminoglycan attachment site:
433-436

N-myristoylation site:
178-183, 181-186, 182-187, 198-203, 339-344, 434-439

Amidation site:
1-4

alpha/beta hydrolase fold:
115-372
```

FIGURE 87

GGCGGCGTCCGTGAGGGGCTCCTTTGGGCAGGGGTAGTGTTTGGTGTCCCTGTCTTGCGT GATATTGACAAACTGAAGCTTTCCTGCACCACTGGACTTAAGGAAGAGTGTACTCGTAGG CGGACAGCTTTAGTGGCCGGCCGGCCGCTCTCATCCCCCGTAAGGAGCAGAGTCCTTTGT ACTGACCAAGATGAGCAACATCTACATCCAGGAGCCTCCCACGAATGGGAAGGTTTTATT GAAAACTACAGCTGGAGATATTGACATAGAGTTGTGGTCCAAAGAAGCTCCTAAAGCTTG CAGAAATTTTATCCAACTTTGTTTGGAAGCTTATTATGACAATACCATTTTTCATAGAGT TGTGCCTGGTTTCATAGTCCAAGGCGGAGATCCTACTGGCACAGGGAGTGGTGGAGAGTC TATCTATGGAGCGCCATTCAAAGATGAATTTCATTCACGGTTGCGTTTTAATCGGAGAGG ACTGGTTGCCATGGCAAATGCTGGTTCTCATGATAATGGCAGCCAGTTTTTCTTCACACT GGGTCGAGCAGATGAACTTAACAATAAGCATACCATCTTTGGAAAGGTTACAGGGGATAC AGTATATAACATGTTGCGACTGTCAGAAGTAGACATTGATGATGACGAAAGACCACATAA TCCACACACACACACACACACCTGTGAGGTTTTGTTTAATCCTTTTGATGACATCATTCCAAG GGAAATTAAAAGGCTGAAAAAAGAGAAACCAGAGGAGGAAGTAAAGAAATTGAAACCCAA AGGCACAAAAATTTTAGTTTACTTTCATTTGGAGAGGAAGCTGAGGAAGAAGAGGAGGA AGTAAATCGAGTTAGTCAGAGCATGAAGGGCAAAAGCAAAAGTAGTCATGACTTGCTTAA GGATGATCCACATCTCAGTTCTGTTCCAGTTGTAGAAAGTGAAAAAGGTGATGCACCAGA TTTAGTTGATGATGAGAGAGATGAAAGTGCAGAGCATGATGAATATATTGATGGTGATGA AAAGAACCTGATGAGAGAAAGAATTGCCAAAAAATTAAAAAAGGACACAAGTGCGAATGT TGCAGCAAAACAAGCAGAAAAAAGAAGTGAAGAGGAAGAAGCCCCTCCAGATGGTGCTGT TGCCGAATACAGAAGAGAAAAGCAAAAGTATGAAGCTTTGAGGAAGCAACAGTCAAAGAA GGGAACTTCCCGGGAAGATCAGACCCTTGCACTGCTGAACCAGTTTAAATCTAAACTCAC TCAAGCAATTGCTGAAACACCTGAAAATGACATTCCTGAAACAGAAGTAGAAGATGATGA AGGATGGATGTCACATGTACTTCAGTTTGAGGATAAAAGCAGAAAAGTGAAAGATGCAAG CATGCAAGACTCAGATACATTTGAAATCTATGATCCTCGGAATCCAGTGAATAAAAGAAG TGATAACCAGAACTTGCTGGAAATGTGCCTACAATGGCCTTGTAACAGCCATTGTTCCCA ACAGCATCACTTAGGGGTGTGAAAAGAAGTATTTTTGAACCTGTTGTCTGGTTTTGAAAA ACAATTATCTTGTTTTGCAAATTGTGGAATGATGTAAGCAAATGCTTTTGGTTACTGGTA CATGTGTTTTTTCCTAGCTGACCTTTTATATTGCTAAATCTGAAATAAAATAACTTTCCT TCCACAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA

FIGURE 88

MSNIYIQEPPTNGKVLLKTTAGDIDIELWSKEAPKACRNFIQLCLEAYYDNTIFHRVVPG FIVQGGDPTGTGSGGESIYGAPFKDEFHSRLRFNRRGLVAMANAGSHDNGSQFFFTLGRA DELNNKHTIFGKVTGDTVYNMLRLSEVDIDDDERPHNPHKIKSCEVLFNPFDDIIPREIK RLKKEKPEEEVKKLKPKGTKNFSLLSFGEEAEBEBEEEVNRVSQSMKGKSKSSHDLLKDDP HLSSVPVVESEKGDAPDLVDDGEDESAEHDEYIDGDEKNLMRERIAKKLKKDTSANVKSA GEGEVEKKSVSRSEELRKEARQLKRELLAAKQKKVENAAKQAEKRSEEEEAPPDGAVAEY RREKQKYEALRKQQSKKGTSREDQTLALLNQFKSKLTQAIAETPENDIPETEVEDDEGWM SHVLOFEDKSRKVKDASMQDSDTFEIYDPRNFVNKRREESKKLMREKKERR

Important features: Signal peptide: amino acids 1-21

N-glycosylation sites: amino acids 109-112 and 201-204

Cyclophilin-type peptidyl-prolyl cis-trans isomerase signature: amino acids 49-66

Homologous region to Cyclophilin-type peptidyl-prolyl cistrans isomerase: amino acids 96-140, 49-89 and 22-51

FIGURE 89

CCCGGCTCCGCTCTGCCCCCTCGGGGTCGCGCCCCACGATGCTGCAGGGCCCTGG $\tt CTCGCTGCTGCTCTTCCTCGCCTCGCACTGCTGCCTGGGCTCGGCGCGCGGGGCTCTT$ CCTCTTTGGCCAGCCCGACTTCTCCTACAAGCGCAGCAATTGCAAGCCCATCCCGGTCAA CCTGCAGCTGTGCCACGGCATCGAATACCAGAACATGCGGCTGCCCAACCTGCTGGGCCA CGAGACCATGAAGGAGGTGCTGGAGCAGGCCGGCGCTTGGATCCCGCTGGTCATGAAGCA GTGCCACCGGACACCAAGAAGTTCCTGTGCTCGCTCTTCGCCCCCGTCTGCCTCGATGA CCTAGACGAGACCATCCAGCCATGCCACTCGCTCTGCGTGCAGGTGAAGGACCGCTGCGC CCCGGTCATGTCCGCCTTCGGCTTCCCCTGGCCCGACATGCTTGAGTGCGACCGTTTCCC GGAAGCTCCAAAGGTATGTGAAGCCTGCAAAAATAAAAATGATGATGACAACGACATAAT GGAAACGCTTTGTAAAAATGATTTTGCACTGAAAATAAAAGTGAAGGAGATAACCTACAT CAACCGAGATACCAAAATCATCCTGGAGACCAAGAGCAAGACCATTTACAAGCTGAACGG TGTGTCCGAAAGGGACCTGAAGAAATCGGTGCTGTGGCTCAAAGACAGCTTGCAGTGCAC CTGTGAGGAGATGAACGACATCAACGCGCCCTATCTGGTCATGGGACAGAAACAGGGTGG $\tt CTCCGCAGCATCCGCAAGCTGCAGTGC\underline{TAG} TCCCGGCATCCTGATGGCTCCGACAGGCC$ TGCTCCAGAGCACGGCTGACCATTTCTGCTCCGGGATCTCAGCTCCCGTTCCCCAAGCAC ACTCCTAGCTGCTCCAGTCTCAGCCTGGGCAGCTTCCCCCTGCCTTTTGCACGTTTGCAT CCCCAGCATTTCCTGAGTTATAAGGCCACAGGAGTGGATAGCTGTTTTCACCTAAAGGAA AAGCCCACCCGAATCTTGTAGAAATATTCAAACTAATAAAATCATGAATATTTTAA

FIGURE 90

MLQGPGSLLLLFLASHCCLGSARGLFLFGQPDFSYKRSNCKPIPVNLQLCHGIEYQNMRL PNLLGHETMKEVLEQAGAWIPLVMKQCHPDTKKFLCSLFAPVCLDDLDETIQPCHSLCVQ VKDRCAPVMSAFGFPWPDMLECDRFPQDNDLCIPLASSDHLLPATEEAPKVCEACKNKND DDNDIMETLCKNDFALKIKVKEITYINRDTKIILETKSKTIYKLNGVSERDLKKSVLWLK DSLQCTCEEMNDINAPYLVMGQRQGGELVITSVKRWQKGQREFKRISRSIRKLQC

Important features:
Signal peptide:
amino acids 1-20

Cysteine rich domain, homolgous to frizzled N terminus: amino acids 6-153

FIGURE 91

GGAAGGGGAGGAGCAGGCCACACAGGCACAGGCCGGTGAGGGACCTGCCCAGACCTGGAG GGTCTCGCTCTGTCACACAGGCTGGAGTGCAGTGGTGTGATCTTGGCTCATCGTAACCTC CACCTCCCGGGTTCAAGTGATTCTCATGCCTCAGCCTCCCGAGTAGCTGGGATTACAGGT GGTGACTTCCAAGAGTGACTCCGTCGGAGGAAAATGACTCCCCAGTCGCTGCTGCAGACG ACACTGTTCCTGCTGAGTCTGCTCTTCCTGGTCCAAGGTGCCCACGGCAGGGGCCACAGG GAAGACTTTCGCTTCTGCAGCCAGCGGAACCAGACACACAGGAGCAGCCTCCACTACAAA CCCACACCAGACCTGCGCATCTCCATCGAGAACTCCGAAGAGGCCCTCACAGTCCATGCC CCTTTCCCTGCAGCCCACCCTGCTTCCCGATCCTTCCCTGACCCCAGGGGCCTCTACCAC TTCTGCCTCTACTGGAACCGACATGCTGGGAGATTACATCTTCTCTATGGCAAGCGTGAC TTCTTGCTGAGTGACAAAGCCTCTAGCCTCCTCTGCTTCCAGCACCAGGAGGAGAGCCTG GCTCAGGGCCCCCCGCTGTTAGCCACTTCTGTCACCTCCTGGTGGAGCCCTCAGAACATC AGCCTGCCCAGTGCCGCCAGCTTCACCTTCTCCTTCCACAGTCCTCCCCACACGGCCGCT CACAATGCCTCGGTGGACATGTGCGAGCTCAAAAGGGACCTCCAGCTGCTCAGCCAGTTC CAGAGCCTGGAGTCGAAACTGACCTCTGTGAGATTCATGGGGGACATGGTGTCCTTCGAG GAGGACCGGATCAACGCCACGGTGTGGAAGCTCCAGCCCACAGCCGGCCTCCAGGACCTG CACATCCACTCCCGGCAGGAGGAGGAGCAGAGCGAGATCATGGAGTACTCGGTGCTGCTG CTGGTGGACTTCAGCAGCCAAGCCCTGTTCCAGGACAAGAATTCCAGCCAAGTCCTGGGT GAGAAGGTCTTGGGGATTGTGGTACAGAACACCAAAGTAGCCAACCTCACGGAGCCCGTG GTGCTCACTTTCCAGCACCAGCTACAGCCGAAGAATGTGACTCTGCAATGTGTTCTGG GTTGAAGACCCCACATTGAGCAGCCCGGGGCATTGGAGCAGTGCTGGGTGTGAGACCGTC AGGAGAGAACCCAAACATCCTGCTTCTGCAACCACTTGACCTACTTTGCAGTGCTGATG GTCTCCTCGGTGGAGGTGGACGCCGTGCACAAGCACTACCTGAGCCTCCTCTCCTACGTG GGCTGTGTCGTCTCTGCCCTGGCCTGCCTTGTCACCATTGCCGCCTACCTCTGCTCCAGG GTGCCCCTGCCGTGCAGGAGGAAACCTCGGGACTACACCATCAAGGTGCACATGAACCTG CTGCTGGCCGTCTTCCTGCTGGACACGAGCTTCCTGCTCAGCGAGCCGGTGGCCCTGACA GGCTCTGAGGCTGCCGAGCCAGTGCCATCTTCCTGCACTTCTCCCTGCTCACCTGC $\tt CTTTCCTGGATGGGCCTCGAGGGGTACAACCTCTACCGACTCGTGGTGGAGGTCTTTGGC$ ACCTATGTCCCTGGCTACCTACTCAAGCTGAGCGCCATGGGCTGGGGCTTCCCCATCTTT CTGGTGACGCTGGTGGCCCTGGTGGATGTGGACAACTATGGCCCCATCATCTTGGCTGTG CATAGGACTCCAGAGGGCGTCATCTACCCTTCCATGTGCTGGATCCGGGACTCCCTGGTC AGCTACATCACCAACCTGGGCCTCTTCAGCCTGGTGTTTCTGTTCAACATGGCCATGCTA GCCACCATGGTGGTGCAGATCCTGCGGCTGCGCCCCCACACCCAAAAGTGGTCACATGTG TTTGCTTCTGGCACCTTCCAGCTTGTCGTCCTCTACCTTTTCAGCATCATCACCTCCTTC CAAGGCTTCCTCATCTTCATCTGGTACTGGTCCATGCGGCTGCAGGCCCGGGGTGGCCCC TCCCCTCTGAAGAGCAACTCAGACAGCGCCAGGCTCCCCATCAGCTCGGGCAGCACCTCG TCCAGCCGCATCTAGGCCTCCAGCCCACCTGCCCATGTGATGAAGCAGAGATGCGGCCTC GTCGCACACTGCCTGTGGCCCCGAGCCAGGCCAGCCCCAGGCCAGTCAGCCGCAGACT AAGTGCGCCGCCATGCTGCCTAGGGTACTGTCCCCACATCTGTCCCAACCCAGCTGGAGG CCTGGTCTCTCCTTACAACCCCTGGGCCCAGCCCTCATTGCTGGGGGCCAGGCCTTGGAT GTTGCTCTGTCTCGTGGTCACCCTGAGGGCACTCTGCATCCTCTGTCATTTTAACCTC AGGTGGCACCCAGGGCGAATGGGGCCCAGGGCAGACCTTCAGGGCCAGAGCCCTGGCGGA

GGAGAGGCCCTTTGCCAGGAGCACAGCAGCAGCTCGCCTACCTCTGAGCCCAGGCCCCCT CCCTCCCTCAGCCCCCAGTCCTCCCTCCATCTTCCCTGGGGTTCTCCTCCTCCCAGG GCCTCCTTGCTCCTTCGTTCACAGCTGGGGGTCCCCGATTCCAATGCTGTTTTTTGGGGA GTGGTTTCCAGGAGCTGCCTGGTGTCTGCTGTAAATGTTTGTCTACTGCACAAGCCTCGG CCTGCCCCTGAGCCAGGCTCGGTACCGATGCGTGGGCTGGGCTAGGTCCCTCTGTCCATC TGGGCCTTTGTATGAGCTGCATTGCCCTTGCTCACCCTGACCAAGCACACGCCTCAGAGG GGCCTCAGCCTCTCAGGCCCTCTTGTGGCAAGAACTGTGGACCATGCCAGTCCCGT CTGGTTTCCATCCACCACTCCAAGGACTGAGACTGACCTCCTCTGGTGACACTGGCCTA GAGCCTGACACTCTCCTAAGAGGTTCTCTCCAAGCCCCCAAATAGCTCCAGGCGCCCTCG GCCGCCCATCATGGTTAATTCTGTCCAACAACACACACGGGTAGATTGCTGGCCTGTTG TAGGTGGTAGGGACACAGATGACCGACCTGGTCACTCCTCCTGCCAACATTCAGTCTGGT ATGTGAGGCGTGCGTGAAGCAAGAACTCCTGGAGCTACAGGGACAGGGAGCCATCATTCC TGCCTGGGAATCCTGGAAGACTTCCTGCAGGAGTCAGCGTTCAATCTTGACCTTGAAGAT GGGAAGGATGTTCTTTTACGTACCAATTCTTTTGTCTTTTGATATTAAAAAGAAGTACA TGTTCATTGTAGAGAATTTGGAAACTGTAGAAGAGAATCAAGAAGAAAAATAAAAATCAG

FIGURE 92

MTPQSLLQTTLFLLSLLFLVQGAHGRGHREDFRFCSQRNQTHRSSLHYKPTPDLRISIEN SEEALTVHAPFPAAHPASRSFPDPRGLYHFCLYWNRHAGRLHLLYGKRDFTLLSDKASSLL CFQHQEESLAQGPPLLATSVTSWWSPQNISLDSAASFTFSFHSPBPPTTAAHNASVDMCELK RDLQLLSQFLKHPQKASRRPSAAPASQQLQSLESKLTSVRFMGDMVSFEEDRINATVWKL QPTAGLQDLHIHSRQEEEQSEIMEYSVLLPRTLFQRTKGRSGEAEKRLLLVDFSSQALFG DKNSSQVLGEKVLGIVVQNTKVANLTEPVULTFQHQLQPKNVTLQCVFWVEDPTLSSPGH WSSAGCETVRRETQTSCFCNHLTYFAVLMVSSVEVDAVHKHYLSLLSYVGCVVSALACLV TIAAYLCSRVPLDCRRKPRDYTIKVHMNLLLAVFLLDTSFLLSEPVALTGSEAGCRASAI FLHFSLLTCLSWMGLEGYNLYRLVVEVFGTYVPGYLLKLKSAMGWGPFIFLTVIVALVDVD NYGPIILAVHRTPEGVIYPSMCWIRDSLVSYITNLGLFSLVFLFNMAMLATMVVQILRLR PHTQKWSRVLTLLGLSLVLGLBWALIFSFSFASGTFQLVVLYFSITTSFQGFLIFIWYWS MRLOARGGPSPLKSNSDSAKLPISSGSTSSRI

Important features:
Signal peptide:
amino acids 1-25

Putative transmembrane domains: amino acida 382-398, 402-420, 445-468, 473-491, 519-537, 568-590 and 634-657

Microbodies C-terminal targeting signal: amino acids 691-693

cAMP- and cGMP-dependent protein kinase phosphorylation sites:

amino acids 198-201 and 370-373

N-glycosylation sites: amino acids 39-42, 148-151, 171-174, 234-237, 303-306, 324-327 and 341-344

G-protein coupled receptors family 2 proteins: amino acids 475 - 504

FIGURE 93

FIGURE 94

 $\label{local_matrix} $$ MKLMVLVFTIGLTLLLGVQAMPANRLSCYRKILKDHNCHNLPEGVADLTQIDVNVQDHFW DGKGCEMICYCNFSELLCCPKDVFFGPKISFVIPCNNQ $$$

Important features: Signal peptide: amino acids 1-20

N-glycosylation site: amino acids 72-76

Tyrosine kinase phosphorylation site: amino acids 63-71

FIGURE 95

FIGURE 96

MKTTTCSLLICISLLQLMVPVNTDETIEIIVENKVKELLANPANYPSTVTKTLSCTSVK TMNRWASCPAGMTATGCACGFACGSWEIQSGDTCNCLCLLVDWTTARCCQLS

FIGURE 97

GAGGCAGAAAGGCAGAAAGGAGAAAATTCAGGATAACTCTCCTGAGGGGTGAGCCAAGCC CTGCCATGTAGTGCACGCAGGACATCAACAAACAGATAACAGGAAATGATCCATTCCC TGTGGTCACTTATTCTAAAGGCCCCAACCTTCAAAGTTCAAGTAGTGATATGGATGACTC GAAGGAGTGTGTTTCCATCCTCCCACGGAAGGAAAGCCCCTCTGTCCGATCCTCCAAAGA CGGAAAGCTGCTGGCAACCTTGCTGCTGGCACTGCTGTCTTGCTGCCTCACGGTGGT GTCTTTCTACCAGGTGGCCGCCCTGCAAGGGGACCTGGCCAGCCTCCGGGCAGAGCTGCA AGCTCCAGCTGTCACCGCGGGACTGAAAATCTTTGAACCACCAGCTCCAGGAGAAGGCAA CTCCAGTCAGAACAGCAGAAATAAGCGTGCCGTTCAGGGTCCAGAAGAAACAGTCACTCA AGACTGCTTGCAACTGATTGCAGACAGTGAAACACCAACTATACAAAAAGGATCTTACAC ATTTGTTCCATGGCTTCTCAGCTTTAAAAGGGGAAGTGCCCTAGAAGAAAAAGAGAATAA AATATTGGTCAAAGAAACTGGTTACTTTTTTATATATGGTCAGGTTTTATATACTGATAA GACCTACGCCATGGGACATCTAATTCAGAGGAAGAAGGTCCATGTCTTTGGGGATGAATT GAGTCTGGTGACTTTGTTTCGATGTATTCAAAATATGCCTGAAACACTACCCAATAATTC CTGCTATTCAGCTGGCATTGCAAAACTGGAAGAAGGAGATGAACTCCAACTTGCAATACC AAGAGAAAATGCACAAATATCACTGGATGGAGATGTCACATTTTTTGGTGCATTGAAACT GCTGTGACCTACTTACACCATGTCTGTAGCTATTTTCCTCCCTTTCTCTGTACCTCTAAG

FIGURE 98

MDDSTEREQSRLTSCLKKREEMKLKECVSILPRKESPSVRSSKDGKLLAATLLLALLSCC LTVVSFYQVAALQGDLASLRAELQGHHAEKLPAGGAPKAGLEEAPAVTAGLKIFBPPAP GEGNSSQNSRNKRAVQGPEETVTQDCLQLIADSETPTIQKGSYTFVPWLLSFKRGSALEE KENKILVKETGYFFIYGQVLYTDKTYAMGHLIQRKKVHVFGDELSLVTLFRCIQNMPETL PNNSCYSAGIAKLEEGDELQLAIPRENAQISLDGDVTFFGALKLL

Transmembrane domain: amino acids 47-72

N-glycosylation site: amino acids 124-127, 242-245

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 33-36, 173-176

N-myristoylation site: amino acids 96-101

TNF family proteins: amino acids 172-206

FIGURE 99

GCGAGGTGGCGATCGCTGAGAGGCAGGAGGGCCGAGGCGGGCCTGGGAGGCGGCCCGGAG GTGGGGCGCCGCTGGGGCCGGCCCGCACGGCTTCATCTGAGGGCGCACGGCCCGCGACC GAGCGTGCGGACTGGCCTCCCAAGCGTGGGGCGACAAGCTGCCGGAGCTGCAAATGGGCCG CGGCTGGGGATTCTTGTTTGGCCTCCTGGGCGCCGTGTGGCTGCTCAGCTCGGGCCACGG AGAGGAGCAGCCCCCGGAGACAGCGGCACAGAGGTGCTTCTGCCAGGTTAGTGGTTACTT GGATGATTGTACCTGTGATGTTGAAACCATTGATAGATTTAATAACTACAGGCTTTTCCC A AGACTACA A A A ACTTCTTGA A AGTGACTACTTTAGGTATTACAAGGTAAACCTGA AGAG GCCGTGTCCTTTCTGGAATGACATCAGCCAGTGTGGAAGAAGGGACTGTGCTGTCAAACC ATGTCAATCTGATGAAGTTCCTGATGGAATTAAATCTGCGAGCTACAAGTATTCTGAAGA AGCCAATAATCTCATTGAAGAATGTGAACAAGCTGAACGACTTGGAGCAGTGGATGAATC TCTGAGTGAGGAAACACAGAAGGCTGTTCTTCAGTGGACCAAGCATGATGATTCTTCAGA TAACTTCTGTGAAGCTGATGACATTCAGTCCCCTGAAGCTGAATATGTAGATTTGCTTCT TAATCCTGAGCGCTACACTGGTTACAAGGGACCAGATGCTTGGAAAATATGGAATGTCAT CTACGAAGAAAACTGTTTTAAGCCACAGACAATTAAAAGACCTTTAAATCCTTTGGCTTC TGGTCAAGGGACAAGTGAAGAGAACACTTTTTACAGTTGGCTAGAAGGTCTCTGTGTAGA AAAAAGAGCATTCTACAGACTTATATCTGGCCTACATGCAAGCATTAATGTGCATTTGAG TGCAAGATATCTTTTACAAGAGACCTGGTTAGAAAAGAAATGGGGACACAACATTACAGA ATTTCAACAGCGATTTGATGGAATTTTGACTGAAGGAGAAGGTCCAAGAAGGCTTAAGAA CTTGTATTTTCTCTACTTAATAGAACTAAGGGCTTTATCCAAAGTGTTACCATTCTTCGA GCGCCAGATTTTCAACTCTTTACTGGAAATAAAATTCAGGATGAGGAAAACAAAATGTT ACTTCTGGAAATACTTCATGAAATCAAGTCATTTCCTTTGCATTTTGATGAGAATTCATT TTTTGCTGGGGATAAAAAAGAAGCACACAAACTAAAGGAGGACTTTCGACTGCATTTTAG AAATATTTCAAGAATTATGGATTGTGTTGGTTGTTTAAATGTCGTCTGTGGGGAAAGCT TCAGACTCAGGGTTTGGGCACTGCTCTGAAGATCTTATTTTCTGAGAAATTGATAGCAAA TATGCCAGAAAGTGGACCTAGTTATGAATTCCATCTAACCAGACAAGAAATAGTATCATT ATTCAACGCATTTGGAAGAATTTCTACAAGTGTGAAAGAATTAGAAAACTTCAGGAACTT GTTACAGAATATTCATTAAAGAAAACAAGCTGATATGTGCCTGTTTCTGGACAATGGAGG CGAAAGAGTGGAATTTCATTCAAAGGCATAATAGCAATGACAGTCTTAAGCCAAACATTT TATATAAAGTTGCTTTTGTAAAGGAGAATTATATTGTTTTAAGTAAACACATTTTTAAAA ATTGTGTTAAGTCTATGTATAATACTACTGTGAGTAAAAGTAATACTTTAATAATGTGGT AAAAAAAAAAAAAAAAAAAAAAAA

FIGURE 100

MGRGWGFLFGLLGAVWLLSSGHGEEQPPETAAQRCFCQVSGYLDDCTCDVETIDRFNNYR LFPRLQKLLESDYFRYYKVNLKRPCPFWNDISQCGRRDCAVKPCQSDEVPDGIKSASYKY SEBANNLIEECEQAERLGAVDESLSEETQKAVLQWTKHDDSSDNFCEADDIQSPEAEYYD LLLMPERYTGYKGPDAWKIWNVIYEENCFKPQTIKRPLNPLASGQGTSEENTFYSWLEGL CVEKRAFYRLISGLHASINVHLSARYLLQETWLEKKWGHNITEFQQRFDGILTEGEGPRR LKNLYFLYNIELRALSKVLPFFRRPDFQLFTGNKIQDEENKWLLLEILHEIKSFPLHFDE NSFFAGDKKEAHKLKEDFRLHFRNISRIMDCVGCFKCRLWGKLQTQGLGTALKILFSEKL IANMPESGPSYEFHLTRQBIVSLFNAFGRISTSVKELENFRNLLQNIH

Important features: Signal peptide: amino acids 1-23

N-glycosylation site: amino acids 280-283 and 384-387

Amidation site: amino acids 94-97

Glycosaminoglycan attachment site: amino acids 20-23 and 223-226

Aminotransferases class-V pyridoxal-phosphate: amino acids 216-222

Interleukin-7 proteins:
amino acids 338-343

FIGURE 101

GCCTAGCCAGGCCAAGAATGCAATTGCCCCGGTGGTGGGAGCTGGGAGACCCCTGTGCTT GGACGGGACAGGGTCGGGGGACACGCAGGATGAGCCCCGCGACCACTGGCACATTCTTGC TGACAGTGTACAGTATTTTCTCCAAGGTACACTCCGATCGGAATGTATACCCATCAGCAG GTGTCCTCTTTGTTCATGTTTTGGAAAGAGAATATTTTAAGGGGGAATTTCCACCTTACC CAAAACCTGGCGAGATTAGTAATGATCCCATAACATTTAATACAAATTTAATGGGTTACC CAGACCGACCTGGATGGCTTCGATATATCCAAAGGACACCATATAGTGATGGAGTCCTAT ATGGGTCCCCAACAGCTGAAAATGTGGGGAAGCCAACAATCATTGAGATAACTGCCTACA ACTTCCCGTTGCCATATCAAGCAGAATTCTTCATTAAGAATATGAATGTAGAAGAAATGT TGGCCAGTGAGGTTCTTGGAGACTTTCTTGGCGCAGTGAAAAATGTGTGGCAGCCAGAGC GCCTGAACGCCATAAACATCACATCGGCCCTAGACAGGGGTGGCAGGGTGCCACTTCCCA TTAATGACCTGAAGGAGGGCGTTTATGTCATGGTTGGTGCAGATGTCCCGTTTTCTTCTT GTTTACGAGAAGTTGAAAATCCACAGAATCAATTGAGATGTAGTCAAGAAATGGAGCCTG TAATAACATGTGATAAAAAATTTCGTACTCAATTTTACATTGACTGGTGCAAAATTTCAT TGGTTGATAAAACAAAGCAAGTGTCCACCTATCAGGAAGTGATTCGTGGAGAGGGGATTT TACCTGATGGTGGAGAATACAAACCCCCTTCTGATTCTTTGAAAAGCAGAGACTATTACA TTGCTTATATCATGTGCTGCCGACGGGAAGGCGTGGAAAAGAGAAACATGCAAACACCAG ACATCCAACTGGTCCATCACAGTGCTATTCAGAAATCTACCAAGGAGCTTCGAGACATGT CCAAGAATAGAGAGATAGCATGGCCCCTGTCAACGCTTCCTGTGTTCCACCCTGTGACTG GGGAAATCATACCTCCTTTACACACAGACAACTATGATAGCACAAACATGCCATTGATGC AAACGCAGCAGAACTTGCCACATCAGACTCAGATTCCCCAACAGCAGACTACAGGTAAAT GGTATCCCTGAAGAAAAAAACTGACTGAAGCAATGAATTTATAATCAGACAATATAGCA GTTACATCACATTTCTTTTCTCTTCCAATAATGCATGAGCTTTTCTGGCATATGTTATGC ATGTTGGCAGTATTAAGTGTATACCAAATAATACAACATAACTTTCATTTTACTAATGTA TTTTTTTTTTTAAAGCATTTTTGACAATTTGTAAAACATTGATGACTTTATATTTTGTT

FIGURE 102

MQLPRWWELGDPCAWTGQGRGTRRMSPATTGTFLLTVYSIFSKVHSDRNVYPSAGVLFVH VLEREYFKGEFPPYPKPGGISNDPITFNTNLMGYPDRPGWLRYIQRTPYSDGVLYGSPTA ENVGKPTIIEITAYNRRTFETARHNLIINIMSAEDFFLPYQAEFFIKMMNVEEMLASEVL GDFLGAVKNVWQPERLNAINITSALDRGGRVPLPINDLKEGYYVMVGADVPFSSCLREVE NPQNQLRCSQEMEPVITCDKKFRTQFYIDWCKISLVDKTKQVSTYQEVIRGEGILPDGGE YKPPSDSLKSRDYYTDFLITLAVPSAVALVLFLILAYIMCCRREGVEKRNMQTPDIQLVH HSAIQKSTKELRDMSKNREIAWPLSTLPVFHPVTGEIIPPLHTDNYDSTNMPLMQTQQNL PHOTOIPQOQTTGKWYP

signal sequence:
Amino acids 1-46

transmembrane domain: Amino acids 319-338

N-glycosylation site: Amino acids 200-204

 $\mathtt{cAMP-}$ and $\mathtt{cGMP-} \mathtt{dependent}$ protein kinase phosphorylation site:

Amino acids 23-27

Tyrosine kinase phosphorylation site: Amino acids 43-52

N-myristoylation sites: Amino acids 17-23;112-118;116-122;185-191

FIGURE 103

CAGAAGAGGGGGCTAGCTGTCTCTGCGGACCAGGGAGACCCCGGGCCCCCCGGT GTGAGGCGGCCTCACAGGGCCGGGTGGGCTGGCGAGCCGACGCGGCGGCGGAGGAGGCTG TGAGGAGTGTGTGGAACAGGACCCGGGACAGAGGAACCATGGCTCCGCAGAACCTGAGCA CCTTTTGCCTGTTGCTGCTATACCTCATCGGGGCGGTGATTGCCGGACGAGATTTCTATA AGATCTTGGGGGTGCCTCGAAGTGCCTCTATAAAGGATATTAAAAAGGCCTATAGGAAAC TAGCCCTGCAGCTTCATCCCGACCGGAACCCTGATGATCCACAAGCCCAGGAGAAATTCC AGGATCTGGGTGCTGCTTATGAGGTTCTGTCAGATAGTGAGAAACGGAAACAGTACGATA CTTATGGTGAAGAAGGATTAAAAGATGGTCATCAGAGCTCCCATGGAGACATTTTTTCAC ACTTCTTTGGGGATTTTGGTTTCATGTTTGGAGGAACCCCTCGTCAGCAAGACAGAAATA TTCCAAGAGGAAGTGATATTATTGTAGATCTAGAAGTCACTTTGGAAGAAGTATATGCAG AGTGCAATTGTCGGCAAGAGATGCGGACCACCCAGCTGGGCCCTGGGCGCTTCCAAATGA TGGAAGTAGAAATAGAGCCTGGGGTGAGAGACGGCATGGAGTACCCCTTTATTGGAGAAG GTGAGCCTCACGTGGATGGGGAGCCTGGAGATTTACGGTTCCGAATCAAAGTTGTCAAGC ACCCAATATTTGAAAGGAGAGAGATGATTTGTACACAAATGTGACAATCTCATTAGTTG AGTCACTGGTTGGCTTTGAGATGGATATTACTCACTTGGATGGTCACAAGGTACATATTT CCCGGGATAAGATCACCAGGCCAGGAGCGAAGCTATGGAAGAAAGGGGAAGGGCTCCCCA ACTTTGACAACAACAATATCAAGGGCTCTTTGATAATCACTTTTGATGTGGATTTTCCAA AAGAACAGTTAACAGAGGAAGCGAGAGAAGGTATCAAACAGCTACTGAAACAAGGGTCAG TGCAGAAGGTATACAATGGACTGCAAGGATAT**TGA**GAGTGAATAAAATTGGACTTTGTTT TTATTTCAATATGCAAGTTAGGCTTAATTTTTTTTTTATCTAATGATCATCATGAAATGAAT AAGAGGGCTTAAGAATTTGTCCATTTGCATTCGGAAAAGAATGACCAGCAAAAGGTTTAC TAATACCTCTCCCTTTGGGGATTTAATGTCTGGTGCTGCCGCCTGAGTTTCAAGAATTAA AGCTGCAAGAGGACTCCAGGAGCAAAAGAAACACAATATAGAGGGTTGGAGTTGTTAGCA ATTTCATTCAAAATGCCAACTGGAGAAGTCTGTTTTTAAATACATTTTGTTGTTATTTTTA

FIGURE 104

MAPQNLSTFCLLLYLIGAVIAGRDFYKILGVPRSASIKDIKKAYRKLALQLHPDRNPDD
PQAQEKFQDLGAAYEVLSDSEKRKQYDTYGEEGLKDGHQSSHGDIFSHFGDFGFMFGGT
PRQQDRNIPRGSDIIVDLEVTLEEVYAGNFVEVVRNKPVARQAPGKRKCNCRQEMRTTQL
GPGRFQMTQEVVCDECPNVKLVNEERTLEVEIEPGVRDGMEYPFIGEGEPHVDGEPGDLR
FRIKVVKHPIFERRGDDLYTNVTISLVBSLVGFEMDITHLDGHKVHLSRDKITRPGAKLW
KKGEGLPNFDNNNIKGSLIIFPDVDFPKEOLTEEAREGIKQLLKQGSVQKVYNGLQGY

Important features:
Signal peptide:
amino acids 1-22

Cell attachment sequence: amino acids 254-257

Nt-dnaJ domain signature: amino acids 67-87

Homologous region to Nt-dnaJ domain proteins:

N-glycosylation site: amino acids 5-9, 261-265

Tyrosine kinase phosphorylation site: amino acids 253-260

N-myristoylation site: amino acids 18-24, 31-37, 93-99, 215-221

Amidation site: amino acids 164-168

FIGURE 105

GGCACGAGGCGGGGCAGTCGCGGGATGCGCCCGGGAGCCACAGCCTGAGGCCCTCAG GTCTCTGCAGGTGTCGTGGAGGAACCTAGCACCTGCCATCCTCTTCCCCAATTTGCCACT TCCAGCAGCTTTAGCCCATGAGGAGGATGTGACCGGGACTGAGTCAGGAGCCCTCTGGAA GCATGGAGACTGTGGTGATTGTTGCCATAGGTGTGCTGGCCACCATCTTTCTGGCTTCGT GCTATGATTCTAAGCCCATTGTGGACCTCATTGGTGCCATGGAGACCCAGTCTGAGCCCT CTGAGTTAGAACTGGACGATGTCGTTATCACCAACCCCCACATTGAGGCCATTCTGGAGA ATGAAGACTGGATCGAAGATGCCTCGGGTCTCATGTCCCACTGCATTGCCATCTTGAAGA TTTGTCACACTCTGACAGAGAGCTTGTTGCCATGACAATGGGCTCTGGGGCCAAGATGA AGACTTCAGCCAGTGTCAGCGACATCATTGTGGTGGCCAAGCGGATCAGCCCCAGGGTGG ATGATGTTGTGAAGTCGATGTACCCTCCGTTGGACCCCAAACTCCTGGACGCACGGACGA CGGGAGGCCTGGACTGGATTGACCAGTCTCTGTCGGCTGCTGAGGAGCATTTGGAAGTCC TTCGAGAACCACCCTACCTTCTGAGCCAGATAAAGGCCTCCCAGGCCCTGAAGGCTTCC TGCAGGAGCAGTCTGCAATT<u>TAG</u>TGCCTACAGGCCAGCAGCTAGCCATGAAGGCCCCTGC CACGGCTGGAGAGTTCAGCTGTGTGTGTGCATAGTAAAGCAGGAGATCCCCGTCAGTTTATG CCTCTTTTGCAGTTGCAAACTGTGGCTGGTGAGTGGCAGTCTAATACTACAGTTAGGGGA GCTCACCTAGTGTTTTCAAGAAAATTGAGCCACCGTCTAAGAAATCAAGAGGTTTCACAT TAAAATTAGAATTTCTGGCCTCTCTCGATCGGTCAGAATGTGTGGCAATTCTGATCTGCA TTTTCAGAAGAGGACAATCAATTGAAACTAAGTAGGGGTTTCTTCTTTTGGCAAGACTTG TACTCTCTCACCTGGCCTGTTTCATTTATTTGTATTATCTGCCTGGTCCCTGAGGCGTCT GGGTCTCTCCTCTCCCTTGCAGGTTTGGGTTTGAAGCTGAGGAACTACAAAGTTGATGAT ATACTTATGTTTCCCTCAAAAAAAAAAAAAAAAA

FIGURE 106

METVVIVAIGVLATIFLASFAALVLVCRQRYCRPRDLLQRYDSKPIVDLIGAMETQSEPS ELELDDVVITNPHTEAILENEDWIEDASGLMSHCIAILKICHTLTEKLVAMTMGSGAKMK TSASVSDIIVVAKRISPRVDDVVKSMYPPLDPKLLDARTTALLLSVSHLVLVTRNACHLT GGLDWIDOSLSAAEEHLEVLREAALASEPDKGLPGPEGFLQEQSAI

FIGURE 107

GCTTCATTTCTCCCGACTCAGCTTCCCACCCTGGGCTTTCCGAGGTGCTTTCGCCGCTGT CCCCACCACTGCAGCCATGATCTCCTTAACGGACACGCAGAAAATTGGAATGGGATTAAC AGGATTTGGAGTGTTTTTCCTGTTCTTTGGAATGATTCTCTTTTTTGACAAAGCACTACT ATTCAGATTCTTCTTCCAAAAACATAAAATGAAAGCTACAGGTTTTTTTCTGGGTGGTGT ATTTGTAGTCCTTATTGGTTGGCCTTTGATAGGCATGATCTTCGAAATTTATGGATTTTT TCTCTTGTTCAGGGGCTTCTTTCCTGTCGTTGTTGGCTTTATTAGAAGAGTGCCAGTCCT TGGATCCCTCCTAAATTTACCTGGAATTAGATCATTTGTAGATAAAGTTGGAGAAAGCAA AGTCATTTGAAGAATATTCAGCACAAAATTAAATTACATGAAATAGCTTGTAATGTTCTT TACAGGAGTTTAAAACGTATAGCCTACAAAGTACCAGCAGCAAATTAGCAAAGAAGCAGT AAATCCATGTTAATGATGCTTAAGAAACTCTTGAAGGCTATTTGTGTTGTTTTTCCACAA TGTGCGAAACTCAGCCATCCTTAGAGAACTGTGGTGCCTGTTTCTTTTCTTTTATTTTG AAGGCTCAGGAGCATCCATAGGCATTTGCTTTTTAGAAGTGTCCACTGCAATGGCAAAAA TATTTCCAGTTGCACTGTATCTCTGGAAGTGATGCATGAATTCGATTGGATTGTCATT CAGGGCC

FIGURE 108

MISLTDTQKIGMGLTGFGVFFLFFGMILFFDKALLAIGNVLFVAGLAFVIGLERTFRFFF QKHKMKATGFFLGGVFVVLIGWPLIGMIFEIYGFFLLFRGFFPVVVGFIRRVPVLGSLLN LPGTRSFVDKVGESNNMV

Important features:
Transmembrane domains:
amino acids 12-30 (typeII), 33-52, 69-89 and 93-109

N-myristoylation sites: amino acids 11-16, 51-56 and 116-121

Aminoacyl-transfer RNA synthetases class-II protein: amino acids 49-59

FIGURE 109

FIGURE 110

MRGTRLALLALVLAÄČGELAPALRCYVCPEPTGVSDCVTIATCTTNETMCKTTLYSREIV YPFOGDSTVTKSCASKCKPSDVDGIGQTLPVSCCNTELCNVDGAPALNSLHCGALTLLPL LSLRL

Important features: Signal peptide: amino acids 1-17

N-glycosylation site: amino acids 46-49

FIGURE 111

GCGCCGCCAGGCGTAGGCGGGGTGGCCCTTGCGTCTCCCGCTTCCTTGAAAAACCCGGCG GGCGAGCGAGGCTGCGGGCCGCCGCCGCCTGCCCTTCCCCACACTCCCCGCCGAGAAGCCTCG CTCGGCGCCCAACATGGCGGGTGGGCGCTGCGGCCCGCAGCTAACGGCGCTCCTGGCCGC CTGGATCGCGGCTGTGGCGGCGACGGCAGGCCCCGAGGAGGCCGCGCTGCCGCGCAGCA GAGCCGGGTCCAGCCCATGACCGCCTCCAACTGGACGCTGGTGATGGAGGGCGAGTGGAT GCTGAAATTTTACGCCCCATGGTGTCCATCCTGCCAGCAGACTGATTCAGAATGGGAGGC TTTTGCAAAGAATGGTGAAATACTTCAGATCAGTGTGGGGAAGGTAGATGTCATTCAAGA ACCAGGTTTGAGTGGCCGCTTCTTTGTCACCACTCTCCCAGCATTTTTTCATGCAAAGGA TGGGATATTCCGCCGTTATCGTGGCCCAGGAATCTTCGAAGACCTGCAGAATTATATCTT GATGTCTGGAATGGCTGGTCTTTTTAGCATCTCTGGCAAGATATGGCATCTTCACAACTA TTTCACAGTGACTCTTGGAATTCCTGCTTGGTGTTCTTATGTGTTTTTCGTCATAGCCAC CTTGGTTTTTGGCCTTTTTATGGGTCTGGTCTTGGTGGTAATATCAGAATGTTTCTATGT GCCACTTCCAAGGCATTTATCTGAGCGTTCTGAGCAGAATCGGAGATCAGAGGAGGCTCA TAGAGCTGAACAGTTGCAGGATGCGGAGGAGGAAAAAGATGATTCAAATGAAGAAGAAAA CAAAGACAGCCTTGTAGATGATGAAGAAGAAGAAGAAGATCTTGGCGATGAGGATGAAGC AGAGGAAGAAGAGGAGGACAACTTGGCTGCTGGTGTGGATGAGGAGAAGTGAGGC CAATGATCAGGGGCCCCCAGGAGAGGACGGTGTGACCCGGGAGGAAGTAGAGCCTGAGGA GGCTGAAGAAGGCATCTCTGAGCAACCCTGCCCAGCTGACACAGAGGTGGTGGAAGACTC $\tt CTTGAGGCAGCGTAAAAGTCAGCATGCTGACAAGGGACTGTAGATTTAATGATGCGTTTT$ CAAGAATACACACCAAAACAATATGTCAGCTTCCCTTTGGCCTGCAGTTTGTACCAAATC CTTAATTTTTCCTGAATGAGCAAGCTTCTCTTAAAAGATGCTCTCTAGTCATTTGGTCTC ATGGCAGTAAGCCTCATGTATACTAAGGAGAGTCTTCCAGGTGTGACAATCAGGATATAG AAAAACAAACGTAGTGTTGGGATCTGTTTGGAGACTGGGATGGGAACAAGTTCATTTACT ${\tt TAGGGGTCAGAGAGTCTCGACCAGAGGAGGCCATTCCCAGTCCTAATCAGCACCTTCCAG}$ AGACAAGGCTGCAGGCCCTGTGAAATGAAAGCCAAGCAGGAGCCTTGGCTCCTGAGCATC AATTGCAGGAAACATCAGGCACCACAGTGCATGAAAAATCTTTCACAGCTAGAAATTGAA AGGGCCTTGGGTATAGAGAGCAGCTCAGAAGTCATCCCAGCCCTCTGAATCTCCTGTGCT ATGTTTTATTTCTTACCTTTAATTTTTCCAGCATTTCCACCATGGGCATTCAGGCTCTCC ACACTCTTCACTATTATCTCTTGGTCAGAGGACTCCAATAACAGCCAGGTTTACATGAAC TGTGTTTGTTCATTCTGACCTAAGGGGTTTAGATAATCAGTAACCATAACCCCTGAAGCT GTGACTGCCAAACATCTCAAATGAAATGTTGTGGCCATCAGAGACTCAAAAGGAAGTAAG GATTTTACAAGACAGATTAAAAAAAAATTGTTTTGTCCAAAATATAGTTGTTGATTT TTTTTTAAGTTTTCTAAGCAATATTTTTCAAGCCAGAAGTCCTCTAAGTCTTGCCAGTAC AAGGTAGTCTTGTGAAGAAAAGTTGAATACTGTTTTGTTTTCATCTCAAGGGGTTCCCTG GGTCTTGAACTACTTTAATAATAACTAAAAAAACCACTTCTGATTTTCCTTCAGTGATGTG CTTTTGGTGAAAGAATTAATGAACTCCAGTACCTGAAAGTGAAAGATTTGATTTTGTTTC CATCTTCTGTAATCTTCCAAAGAATTATATCTTTGTAAATCTCTCAATACTCAATCTACT GTAAGTACCCAGGGAGGCTAATTTCTTT

FIGURE 112

MAGGRCGPQLTALLAAWIAAVAATAGPEEAALPPEQSRVQPMTASNWTLVMEGEWMLKFY APWGPSCQQTDSWEAFAKNGEILQISVGKVDVIQEFGLSGRFFVTTLPAFFHAKDGIF RYRGGPGIFEDLONYILEKKWQSVEPLTGWKSPASLTMSGMAGLFSISGKIWHLHNYFTVT LGIPAWCSYVFFVIATLVFGLFMGLVLVVISECFYVPLPRHLSERSEQNRRSEEAHRAEQ LQDAEEEKDDSNEEENKDSLVDDEEEKEDLGDEDEAEEEEEEDNLAAGVDEERSEANDQG PPGBDGYTREFVEPEEAEEGISEOPCPADTEVVEDSLRORKSOHADKGL

Important features: Signal peptide: amino acids 1-22

Transmembrane domain: amino acids 191-211

N-glycosylation site: amino acids 46-49

Thioredoxin family proteins: (homologous region to disulfide isomerase) amino acids 56-72

Flavodoxin proteins: amino acids 173-187

FIGURE 113

GAGGAACCTACCGGTACCGGCCGCGCGCTGGTAGTCGCCGGTGTGGCTGCACCTCACCAA TCCGTGCGCGGCTGGGCCGTCGGAGAGTGCGTGTGCTTCTCTCCTGCACGCGGTGC TTGGGCTCGGCCAGGCTCCGCCGCCAGGGTTTGAGGATGGGGGAGTAGCTACAGGA AGCGACCCGCGATGGCAAGGTATATTTTTGTGGAATGAAAAGGAAGTATTAGAAATGAG CTGAAGACCATTCACAGATTAATATTTTTGGGGACAGATTTGTGATGCTTGATTCACCCT TGAAGTAATGTAGACAGAAGTTCTCAAATTTGCATATTACATCAACTGGAACCAGCAGTG TAAAGATGACTATATCAGAGACTTGAAAAGGATCATTCTCTGTTTTCTGATAGTGTATAT GGCCATTTTAGTGGGCACAGATCAGGATTTTTACAGTTTACTTGGAGTGTCCAAAACTGC AAGCAGTAGAGAAATAAGACAAGCTTTCAAGAAATTGGCATTGAAGTTACATCCTGATAA A A A C C CGA A TA A C C CA A A TG C A C A TG C G C G A TT TT TT A A A A A TA A A TA G G C G A TA TG A A G C ACTCAAAGATGAAGATCTACGGAAAAAGTATGACAAATATGGAGAAAAGGGACTTGAGGA TAATCAAGGTGGCCAGTATGAAAGCTGGAACTATTATCGTTATGATTTTGGTATTTATGA TGATGATCCTGAAATCATAACATTGGAAAGAAGAGAATTTGATGCTGCTGTTAATTCTGG AGAACTGTGGTTTGTAAATTTTTACTCCCCAGGCTGTTCACACTGCCATGATTTAGCTCC CACATGGAGAGACTTTGCTAAAGAAGTGGATGGGTTACTTCGAATTGGAGCTGTTAACTG TGGTGATGATAGAATGCTTTGCCGAATGAAAGGAGTCAACAGCTATCCCAGTCTCTTCAT TTTTCGGTCTGGAATGGCCCCAGTGAAATATCATGGAGACAGATCAAAGGAGAGTTTAGT GAGTTTTGCAATGCAGCATGTTAGAAGTACAGTGACAGAACTTTGGACAGGAAATTTTGT AGGAGGAGATTGTTTGACTTCACAGACACGACTCAGGCTTAGTGGCATGTTGTTTCTCAA CTCATTGGATGCTAAAGAAATATATTTGGAAGTAATACATAATCTTCCAGATTTTGAACT ACTTTCGGCAAACACACTAGAGGATCGTTTGGCTCATCATCGGTGGCTGTTATTTTTTCA TTTTGGAAAAAATGAAAATTCAAATGATCCTGAGCTGAAAAAACTAAAAACTCTACTTAA AAATGATCATATTCAAGTTGGCAGGTTTGACTGTTCCTCTGCACCAGACATCTGTAGTAA TCTGTATGTTTTCAGCCGTCTCTAGCAGTATTTAAAGGACAAGGAACCAAAGAATATGA TTCTCATGTTACCACGCTTGGACCTCAAAATTTTCCTGCCAATGACAAAGAACCATGGCT TGTTGATTTCTTTGCCCCCTGGTGTCCACCATGTCGAGCTTTACTACCAGAGTTACGAAG AGCATCAAATCTTCTTTATGGTCAGCTTAAGTTTGGTACACTAGATTGTACAGTTCATGA GGGACTCTGTAACATGTATAACATTCAGGCTTATCCAACAACAGTGGTATTCAACCAGTC CAACATTCATGAGTATGAAGGACATCACTCTGCTGAACAAATCTTGGAGTTCATAGAGGA TCTTATGAATCCTTCAGTGGTCTCCCTTACACCCACCACCTTCAACGAACTAGTTACACA AAGAAAACACAACGAAGTCTGGATGGTTGATTTCTATTCTCCGTGGTGTCATCCTTGCCA AGTCTTAATGCCAGAATGGAAAAGAATGGCCCGGACATTAACTGGACTGATCAACGTGGG CAGTATAGATTGCCAACAGTATCATTCTTTTTGTGCCCAGGAAAACGTTCAAAGATACCC TGAGATAAGATTTTTCCCCCAAAATCAAATAAAGCTTATCAGTATCACAGTTACAATGG $\tt TTGGAATAGGGATGCTTATTCCCTGAGAATCTGGGGTCTAGGATTTTTACCTCAAGTATC$ CACAGATCTAACACCTCAGACTTTCAGTGAAAAAGTTCTACAAGGGAAAAATCATTGGGT GATTGATTTCTATGCTCCTTGGTGTGGACCTTGCCAGAATTTTGCTCCAGAATTTGAGCT CTTGGCTAGGATGATTAAAGGAAAAGTGAAAGCTGGAAAAGTAGACTGTCAGGCTTATGC TCAGACATGCCAGAAAGCTGGGATCAGGGCCTATCCAACTGTTAAGTTTTATTTCTACGA AAGAGCAAAGAGAAATTTTCAAGAAGAGCAGATAAATACCAGAGATGCAAAAGCAATCGC TGCCTTAATAAGTGAAAAATTGGAAACTCTCCGAAATCAAGGCAAGAGGAATAAGGATGA ACTTTGATAATGTTGAAGATGAAGAAAAAGTTTAAAAGAAATTCTGACAGATGACATCAG

GCAGTTGTACTGCCAGAATTATCTACAGCACTGGTGTAAAAGAAGGGTCTGCAAACTTTT TCTGTAAAGGGCCGGTTTATAAATATTTTAGACTTTGCAGGCTATAATATATGGTTCACA CATGAGAACAAGAATAGAGTCATCATGTATTCTTTGTTATTTGCTTTTAACAACCTTTAA AAAATATTAAAACGATTCTTAGCTCAGAGCCATACAAAAGTAGGCTGGATTCAGTCCATG TGAGTCTGCTGTGCTATCTACATAAATGTCTAAGTTGTATAAAGTCCACTTTCCCTTCAC GTTTTTTGGCTGACCTGAAAAGAGGTAACTTAGTTTTTGGTCACTTGTTCTCCTAAAAAT GCTATCCCTAACCATATATTTATATTTCGTTTTAAAAAACACCCATGATGTGGCACAGTAA AAGGTTGAAAAAATGCTTTTAATTTTTCACAGCCGAGAAACAGTGCAGCAGTATATGTGC ACACAGTAAGTACACAAATTTGAGCAACAGTAAGTGCACAAATTCTGTAGTTTGCTGTAT CATCCAGGAAAACCTGAGGGAAAAAAATTATAGCAATTAACTGGGCATTGTAGAGTATCC TAAATATGTTATCAAGTATTTAGAGTTCTATATTTTAAAGATATATGTGTTCATGTATTT TCTGAAATTGCTTTCATAGAAATTTTCCCACTGATAGTTGATTTTTTGAGGCATCTAATAT TACTTTACAGGTTGTTTTACTGTAGCTTATAATGATACTGTAGTTATTCCAGTTACTAGT TTACTGTCAGAGGGCTGCCTTTTTCAGATAAATATTGACATAATAACTGAAGTTATTTTT ATAAGAAAATCAAGTATAAAATCTAGGAAAGGGATCTTCTAGTTTCTGTGTTTTAGA CTCAAAGAATCACAAATTTGTCAGTAACATGTAGTTGTTTAGTTATAATTCAGAGTGTAC AGAATGGTAAAAATTCCAATCAGTCAAAAGAGGTCAATGAATTAAAAGGCTTGCAACTTT TTCAAAAAAAAAAAAAAAAA

FIGURE 114

MGVWLNKDDYIRDLKRIILCFLIVYMAILVGTDQDFYSLLGVSKTASSREIRQAFKKLAL
KLHPDKNPNNPNAHGDFLKINRAYEVLKDEDLRKKYDKYGEKGLEDNQGGYESWNYYRY
DFGIYDDDPEITTLERREFDAAVNSGELWFVNFYSPGCSCHCHDLAPTWRDFAKEVDGLLR
IGAVNCGDDRMLCRMKGVNSYPSLFIFRSGMAPVKYHGDRSKESLVSFAMQHVRSTVTEL
WTGNFVNSIQTAFAAGIGWLITFCSKGGDCLTSQTRIRLSGMLFLNSLDAKEIYLEVIHN
DLPDFELLSANTLEDRLAHHRWLLFHFFKKNENSNDPBLKKLKTLLKUNDHIQVGRFDCSSA
PDLCSNLTVFQPSLAVFKGQGTKEYEIHHGKKILYDILAFAKESVNSHVTTLGPQNFPAN
DKEBWLVDFFAPWCPPCRALLPELRRASNLLYGQLKFGTLDCTVHEGLCNMYNIQAYPTI
VVFNQSNIHEYEGHHSAEQILEFIEDLMNPSVVSLTPTTFNELVTQRKHNEVWNDFYSP
WCHPCQVLMPEWKRMARTLTGLINVGSIDCQQYHSFCAQENVQRYPEIRFFPPKSNKAYQ
YHSYNGWNRDAYSLRIWGLGFLPQVSTDLTPQTFSEKVLQGKNHWVIDFYAPWCGPCQNF
APEFELLARMIKGKVKAGKVDCQAYAQTCQKAGIRAYPTVKFYFYERAKRNFQEEQINTR
DAKAIAALISEKLETLENQGKRNKDEL

Important features: Endoplasmic reticulum targeting sequence: amino acids 744-747

Cytochrome c family heme-binding site signature: amino acids 158-163

Nt-dnaJ domain signature: amino acids 77-96

N-glycosylation site: amino acids 484-487

FIGURE 115

GCGGGCTGTTGACGGCGCTGCGATGCCTGCCTGCGAGGGCAGAAGCGGAGCTCTCGG TTCCTCTCAGTCGGACTTCCTGACGCCGCCAGTGGGCGGGGCCCCTTGGGCCGTCGCCAC CACTGTAGTCATGTACCCACCGCCGCCGCCGCCGCCTCATCGGGACTTCATCTCGGTGAC GCTGAGCTTTGGCGAGAGCTATGACAACAGCAAGAGTTGGCGGCGCGCTCGTGCTGGAG GAAATGGAAGCAACTGTCGAGATTGCAGCGGAATATGATTCTCTTCCTCCTTGCCTTTCT GCTTTTCTGTGGACTCCTCTTCTACATCAACTTGGCTGACCATTGGAAAGCTCTGGCTTT CAGGCTAGAGGAAGAGCAGAAGATGAGGCCAGAAATTGCTGGGTTAAAACCAGCAAATCC ACCCGTCTTACCAGCTCCTCAGAAGGCGGACACCGACCCTGAGAACTTACCTGAGATTTC GTCACAGAAGACACAAAGACACATCCAGCGGGGACCACCTCACCTGCAGATTAGACCCCC AAGCCAAGACCTGAAGGATGGGACCCAGGAGGAGGCCACAAAAAGGCAAGAAGCCCCTGT GGATCCCCGCCGGAAGGAGATCCGCAGAGGACAGTCATCAGCTGGAGGGGAGCGGTGAT CGAGCCTGAGCAGGGCACCGAGCTCCCTTCAAGAAGAGCAGAAGTGCCCACCAAGCCTCC $\tt CCTGCCACCGGCCAGGACACAGGGCACACCAGTGCATCTGAACTATCGCCAGAAGGGCGT$ GATTGACGTCTTCCTGCATGCATGGAAAGGATACCGCAAGTTTGCATGGGGCCATGACGA GCTGAAGCCTGTGTCCAGGTCCTTCAGTGAGTGGTTTTGGCCTCGGTCTCACACTGATCGA CGCGCTGGACACCATGTGGATCTTGGGTCTGAGGAAAGAATTTGAGGAAGCCAGGAAGTG GGTGTCGAAGAAGTTACACTTTGAAAAGGACGTGGACGTCAACCTGTTTGAGAGCACGAT CCGCATCCTGGGGGGGCTCCTGAGTGCCTACCACCTGTCTGGGGACAGCCTCTTCCTGAG GAAAGCTGAGGATTTTGGAAATCGGCTAATGCCTGCCTTCAGAACACCATCCAAGATTCC TTACTCGGATGTGAACATCGGTACTGGAGTTGCCCACCCGCCACGGTGGACCTCCGACAG CACTGTGGCCGAGGTGACCAGCATTCAGCTGGAGTTCCGGGAGCTCTCCCGTCTCACAGG GGATAAGAAGTTTCAGGAGGCAGTGGAGAAGGTGACACAGCACATCCACGGCCTGTCTGG GGGCGTATTCACGCTGGGCGCCAGGGCCGACAGCTACTATGAGTACCTGCTGAAGCAGTG GATCCAGGGCGGGAAGCAGGAGACACAGCTGCTGGAAGACTACGTGGAAGCCATCGAGGG TGTCAGAACGCACCTGCTGCGGCACTCCGAGCCCAGTAAGCTCACCTTTGTGGGGGAGCT TGCCCACGGCCGCTTCAGTGCCAAGATGGACCACCTGGTGTGCTTCCTGCCAGGGACGCT GGCTCTGGGCGTCTACCACGGCCTGCCCGCCAGCCACATGGAGCTGGCCCAGGAGCTCAT GGAGACTTGTTACCAGATGAACCGGCAGATGGAGACGGGGCTGAGTCCCGAGATCGTGCA CTTCAACCTTTACCCCCAGCCGGGCCGTCGGGACGTGGAGGTCAAGCCAGCAGACAGGCA CAACCTGCTGCGGCCAGAGACCGTGGAGAGCCTGTTCTACCTGTACCGCGTCACAGGGGA CCGCAAATACCAGGACTGGGGCTGGGAGATTCTGCAGAGCTTCAGCCGATTCACACGGGT CCCCTCGGGTGGCTATTCTTCCATCAACAATGTCCAGGATCCTCAGAAGCCCGAGCCTAG GGACAAGATGGAGAGCTTCTTCCTGGGGGAGACGCTCAAGTATCTGTTCTTGCTCTTCTC CGATGACCCAAACCTGCTCAGCCTGGACGCCTACGTGTTCAACACCGAAGCCCACCCTCT $\tt GCCTATCTGGACCCCTGCC\underline{TAG} GGTGGATGGCTGCTGGTGTGGGGACTTCGGGTGGGCAG$ AGGCACCTTGCTGGGTCTGTGGCATTTTCCAAGGGCCCACGTAGCACCGGCAACCGCCAA GTGGCCCAGGCTCTGAACTGGCTCTGGGCTCCTCCTCGTCTCTGCTTTAATCAGGACACC GTGAGGACAAGTGAGGCCGTCAGTCTTGGTGTGATGCGGGGTGGGCTGGGCCGCTGGAGC CTCCGCCTGCTTCCTCCAGAAGACACGAATCATGACTCACGATTGCTGAAGCCTGAGCAG GTCTCTGTGGGCCGACCAGAGGGGGGCTTCGAGGTGGTCCCTGGTACTGGGGTGACCGAG TGGACAGCCCAGGGTGCAGCTCTGCCCGGGCTCGTGAAGCCTCAGATGTCCCCAATCCAA TACAAGCTGGACTCAGGGATCCTCCTGGCCGCCCCGCAGGGGGCTTGGAGGGCTGGACGG CAAGTCCGTCTAGCTCACGGGCCCCTCCAGTGGAATGGGTCTTTTCGGTGGAGATAAAAG TTGATTTGCTCTAACCGCAA

FIGURE 116

MAACEGRRSGALGSSQSDFLTPPVGGAPWAVATTVVMYPPPPPPPPHRDFISVTLSFGESY
DNSKSWRRRSCWRKWKOLSRLORNMILPLLAFLLFCGLLFYINLADHWKALAFRLEEEQK
MRPEIAGLKPANPPVLPAPQKADTDPENLPEISSQKTORHIQRGPPHLQIRPPSQDLKDG
TQEEATKRQEAPVDPRPEGDPQRTVISWRGAVIEPEQGTELPSRRAEVPTKPPLPPARTO
GTPVHLNYRQKGYIDVFLHAWKGYRKFAWGHDELKPVSRSFSEWFGLGLTLIDALDTMWI
LGLRKEFEEARKWVSKKLHFEKDDVNLIFESTIRILGGLLSAYHLSGDSLFLRKAEDFGN
RLMPAFRTPSKIPYSDVNIGTGVAHPPRWTSDSTVAEVTSIQLEFRELSRITGDKKFQEA
VEKVTQHIHGLSGKKDGLVPMFINTHSGLFTHLGVFTLGARADSYYEYLLKQWIQGKQE
TQLLEDYVEAIEGYRTHLLRHSEPSKLTFVGELAHGRFSAKMDHLVCFLPGTLALGVYHG
LPASHMELAQELMETCYQMNRQMETGLSPEIVHFNLYPQPGRRDVEVKPADRHNLLRPET
VESLFYLYRVTGDRKYQDWGWEILQSFSRFTRVPSGGYSSINNVQDPQKPEPDKMESFF
LGETLKYLFLLFSDDPNLLSLDAYVFNTEAHPLPIWTPA

Important features of the protein: Transmembrane domain: amino acids 21-40 and 84-105 (type II)

FIGURE 117

GTGGGATTTATTTGAGTGCAAGATCGTTTTCTCAGTGGTGGTGGAAGTTGCCTCATCGCA GGCAGATGTTGGGGCTTTGTCCGAACAGCTCCCCTCTGCCAGCTTCTGTAGATAAGGGTT AAAAACTAATATTTATATGACAGAAGAAAAG**ATG**TCATTCCGTAAAGTAAACATCATCA TCTTGGTCCTGGCTGTTGCTCTTCTTACTGGTTTTTGCACCATAACTTCCTCAGCTTGA GCAGTTTGTTAAGGAATGAGGTTACAGATTCAGGAATTGTAGGGCCTCAACCTATAGACT TTGTCCCAAATGCTCTCCGACATGCAGTAGATGGGAGACAAGAGGAGATTCCTGTGGTCA TCGCTGCATCTGAAGACAGGCTTGGGGGGGCCATTGCAGCTATAAACAGCATTCAGCACA ACACTCGCTCCAATGTGATTTTCTACATTGTTACTCTCAACAATACAGCAGACCATCTCC CCTCCTCCCTCAACACTCATTCCCTGAAAACCATCAGATACAAAATTGTCAATTTTGACC CTAAACTTTTGGAAGGAAAGTAAAGGAGGATCCTGACCAGGGGGAATCCATGAAACCTT TAACCTTTGCAAGGTTCTACTTGCCAATTCTGGTTCCCAGCGCAAAGAAGCCCATATACA TGGATGATGATGTAATTGTGCAAGGTGATATTCTTGCCCTTTACAATACAGCACTGAAGC CAGGACATGCAGCTGCATTTTCAGAAGATTGTGATTCAGCCTCTACTAAAGTTGTCATCC GTGGAGCAGGAAACCAGTACAATTACATTGGCTATCTTGACTATAAAAAGGAAAGAATTC GTAAGCTTTCCATGAAAGCCAGCACTTGCTCATTTAATCCTGGAGTTTTTGTTGCAAACC TGACGGAATGGAAACGACAGAATATAACTAACCAACTGGAAAAATGGATGAAACTCAATG TAGAAGAGGGACTGTATAGCAGAACCCTGGCTGGTAGCATCACAACACCTCCTCTGCTTA TCGTATTTTATCAACAGCACTCTACCATCGATCCTATGTGGAATGTCCGCCACCTTGGTT CCAGTGCTGGAAAACGATATTCACCTCAGTTTGTAAAGGCTGCCAAGTTACTCCATTGGA ATGGACATTTGAAGCCATGGGGAAGGACTGCTTCATATACTGATGTTTTGGGAAAAATGGT ATATTCCAGACCCAACAGGCAAATTCAACCTAATCCGAAGATATACCGAGATCTCAAACA TAAAGTGAAACAGAATTTGAACTGTAAGCAAGCATTTCTCAGGAAGTCCTGGAAGATAGC ATGCATGGGAAGTAACAGTTGCTAGGCTTCAATGCCTATCGGTAGCAAGCCATGGAAAAA GATGTGTCAGCTAGGTAAAGATGACAAACTGCCCTGTCTGGCAGTCAGCTTCCCAGACAG ACTATAGACTATAAATATGTCTCCATCTGCCTTACCAAGTGTTTTCTTACTACAATGCTG

FIGURE 118

MSFRKVNIIILVLAVALFLLVLHHNFLSLSSLLRNEVTDSGIVGPQPIDFVPNALRHAVD GRQEEIPVVIAASEDRLGGAIAAINSIQHNTRSNVIFYIVTLNNTADHLRSWLNSDSLKS IRYKIVNFDPKLLEGKVKEDPDQGESMKPLTFARFYLPILVPSAKKAIYMDDDVIVQGDI LALYNTALKPGHAAAFSEDCDSASTKVVIRGAGNQYNYIGYLDYKKERIRKLSMKASTCS FNPGVFVANLTEWKRQNITNQLEKWMKLNVEEGLYSRTLAGSITTPPLLIVFYQQHSTID PMWNVRHLGSSAGKRYSPQFVKAAKLHWNGHLKPWGRTASYTDVWEKWYIPDPTGKFNL IRRYTEISNIK

FIGURE 119

FIGURE 120

MLPPMALPSVSWMLLSCLILLCQVQGEETQKELPSPRISCPKGSKAYGSPCYALFLSPKS WMDADLACQKRPSGKLVSVLSGAEGSFVSSLVRSISNSYSYIWIGLHDPTQGSEPDGDGW EWSSTDVNNYFAWEKNPSTILNPGHCGSLSRSTGFLKWKDYNCDAKLPYVCKFKD

Important features: Signal peptide: amino acids 1-26

C-type lectin domain signature: amino acids 146-171

FIGURE 121

AAAGTTACATTTTCTCTGGAACTCTCCTAGGCCACTCCCTGCTGATGCAACATCTGGGTT TGGGCAGAAAGGAGGTGCTTCGGÁGCCCGCCCTTTCTGAGCTTCCTGGGCCGGCTCTAG AACAATTCAGGCTTCGCTGCGACTCAGACCTCAGCTCCAACATATGCATTCTGAAGAAAG ATGGCTGAGATGGACAGAATGCTTTATTTTGGAAAGAAACAATGTTCTAGGTCAAACTGA GTCTACCAAATGCAGACTTTCACAATGGTTCTAGAAGAAATCTGGACAAGTCTTTTCATG CCTCAGAACCTCTCTGTACTCTCAACCAACATGAAGCATCTCTTGATGTGGAGCCCAGTG ATCGCGCCTGGAGAAACAGTGTACTATTCTGTCGAATACCAGGGGGAGTACGAGAGCCTG GATGTCACTGATGACATCACGGCCACTGTGCCATACAACCTTCGTGTCAGGGCCACATTG GGCTCACAGACCTCAGCCTGGAGCATCCTGAAGCATCCCTTTAATAGAAACTCAACCATC CTTACCCGACCTGGGATGGAGATCACCAAAGATGGCTTCCACCTGGTTATTGAGCTGGAG GAACATGTCAAAATGGTGAGGAGTGGGGGTATTCCAGTGCACCTAGAAACCATGGAGCCA GGGGCTGCATACTGTGAAGGCCCAGACATTCGTGAAGGCCATTGGGAGGTACAGCGCC ${\tt TTCAGCCAGACAGAATGTGTGGAGGTGCAAGGAGAGGCCATTCCCCTGGTACTGGCCCTG}$ TTTGCCTTTGTTGGCTTCATGCTGATCCTTGTGGTCGTGCCACTGTTCGTCTGGAAAATG GGCCGGCTGCTCCAGTACTCCTGTTGCCCCGTGGTGGTCCTCCCAGACACCTTGAAAATA ACCAATTCACCCCAGAAGTTAATCAGCTGCAGAAGGGAGGAGGTGGATGCCTGTGCCACG GCTGTGATGTCTCCTGAGGAACTCCTCAGGGCCTGGATCTCATAGGTTTGCGGAAGGGCC CAGGTGAAGCCGAGAACCTGGTCTGCATGACATGGAAACCATGAGGGGACAAGTTGTGTT TCTGTTTTCCGCCACGGACAAGGGATGAGAGAGTAGGAAGAGCCTGTTGTCTACAAGTC GATGTGACCTCTAGACTGGGGGCTGCCACTTGCTGGCTGAGCAACCCTGGGAAAAGTGAC $\tt TTCATCCCTTCGGTCCTAAGTTTTCTCATCTGTAATGGGGGAATTACCTACACACCTGCT$ AAACACACACACAGAGTCTCTCTCTATATATACACACGTACACATAAATACACCCAGC GAGAGCAGGACATAAATGTATGATGAGAATGATCAAGGACTCTACACACTGGGTGGCTTG GAGAGCCCACTTTCCCAGAATAATCCTTGAGAGAAAAGGAATCATGGGAGCAATGGTGTT GAGTTCACTTCAAGCCCAATGCCGGTGCAGAGGGGAATGGCTTAGCGAGCTCTACAGTAG GTGACCTGGAGGAAGGTCACAGCCACACTGAAAATGGGATGTGCATGAACACGGAGGATC CATGAACTACTGTAAAGTGTTGACAGTGTGTGCACACTGCAGACAGCAGGTGAAATGTAT TTTCTGTTGGTAAAGTACAGAATTCAGCAAATAAAAAGGGCCACCCTGGCCAAAAGCGGT AAAAAAAAAAAAA

FIGURE 122

MQTFTMVLEEIWTSLFMWFFYALIPCLLTDEVAILPAPQNLSVLSTNMKHLLMWSPVIAP GETVYYSVEYQGEYESLYTSHIWIPSSWCSLTEGPECDVTDDITATVPYNLRVRATLGSY TSAWSILKHFPNRNSTILTRPGMEITKDGFHLVIELEDLGPQFEFLVAYWRREPGABEHV KMVRSGGIPVHLETMEPGAAYCVKAQTFVKAIGRYSAFSQTECVEVQGEAIPLVLALFAF VGFMLILVVVPLFVWKMGRLLQYSCCPVVVLPDTLKITNSPQKLISCRREEVDACATAVM SPEEILLRAWIS

Important features: Signal peptide: amino acids 1-29

Transmembrane domain: amino acids 230-255

N-glycosylation sites: amino acids 40-43 and 134-137

Tissue factor proteins homology: amino acids 92-119

Integrins alpha chain protein homology:
amino acids 232-262

FIGURE 123

CGGACGCGTGGGCCGCCACCTCCGGAACAAGCCATGGTGGCGGCGACGGTGGCAGCGGCG TGGCTGCTCCTGTGGGCTGCGCCTGCGCGCAGCAGCAGCAGGACTTCTACGACTTCAAG GCGGTCAACATCCGGGGCAAACTGGTGTCGCTGGAGAAGTACCGCGGATCGGTGTCCCTG GTGGTGAATGTGGCCAGCGAGTGCGGCTTCACAGACCAGCACTACCGAGCCCTGCAGCAG CTGCAGCGAGACCTGGGCCCCCACCACTTTAACGTGCTCGCCTTCCCCTGCAACCAGTTT GGCCAACAGGAGCCTGACAGCAACAAGGAGATTGAGAGCTTTGCCCGCCGCACCTACAGT GTCTCATTCCCCATGTTTAGCAAGATTGCAGTCACCGGTACTGGTGCCCATCCTGCCTTC AAGTACCTGGCCCAGACTTCTGGGAAGGAGCCCACCTGGAACTTCTGGAAGTACCTAGTA GCCCAGATGGAAAGGTGGTAGGGGCTTGGGACCCAACTGTGTCAGTGGAGGAGGTCAGA CCCCAGATCACAGCGCTCGTGAGGAAGCTCATCCTACTGAAGCGAGAAGACTTA**TAA**CCA CCGCGTCTCCTCCACCACCTCATCCCGCCCACCTGTGTGGGGCTGACCAATGCAAAC GGTCCCATCATTCTTGTGGGGGAAAAATTCTAGTATTTTGATTATTTGAATCTTACAGCA ACAAATAGGAACTCCTGGCCAATGAGAGCTCTTGACCAGTGAATCACCAGCCGATACGAA AGGCTTCTGTAAACTGGGACCAATGATTACCTCATAGGGCTGTTGTGAGGATTAGGATGA AATACCTGTGAAAGTGCCTAGGCAGTGCCAGCCAAATAGGAGGCATTCAATGAACATTTT TTGCATATAAACCAAAAAATAACTTGTTATCAATAAAAACTTGCATCCAACATGAATTTC CAGCCGATGATAATCCAGGCCAAAGGTTTAGTTGTTGTTATTTCCTCTGTATTATTTTCT TCATTACAAAGAATGCAAGTTCATTGTAACAATCCAAACAATACCTCACGATATAAAA TAAAAATGAAAGTATCCTCCTCAAAAA

FIGURE 124

MVAATVAAAWLLLWAAACAQQEQDFYDFKAVNIRGKLVSLEKYRGSVSLVVNVASECGFT DQHYRALQQLQRDLGPHHFNVLAFPCNQFGQQEPDSNKEIESFARRTYSVSFPMFSKIAV TGTGAHPAFKYLAQTSGKEPTWNFWKYLVAPDGKVVGAWDPTVSVEEVRPQITALVRKLI LLKREDL

FIGURE 125

CGGACGCGTGGGCGGACGCGTGGGCGGACGCGTGGGTTGGGAGGGGGCAGGATGGGAGGG AAAGTGAAGAAAACAGAAAAGGAGAGGGACAGAGGCCAGAGGACTTCTCATACTGGACAG AAACCGATCAGGCATGGAACTCCCCTTCGTCACTCACCTGTTCTTGCCCCTGGTGTTCCT GACAGGTCTCTGCTCCCCCTTTAACCTGGATGAACATCACCCACGCCTATTCCCAGGGCC ACCAGAAGCTGAATTTGGATACAGTGTCTTACAACATGTTGGGGGTGGACAGCGATGGAT GCTGGTGGGCGCCCCCTGGGATGGGCCTTCAGGCGACCGGAGGGGGGACGTTTATCGCTG CCCTGTAGGGGGGCCCACAATGCCCCATGTGCCAAGGGCCACTTAGGTGACTACCAACT GGGAAATTCATCTCATCCTGCTGTGAATATGCACCTGGGGATGTCTCTGTTAGAGACAGA TGGTGATGGGGGATTCATGGTGAGCTAAGGAGGGTGGTGGCAGTGTCTCTGAAGGTCC ATAAAAGAAAAAGAGAGTGTGGTAAGGGAAAATGGTCTGTGGAGGGGTCAAGGAGT TAAAAACCCTAGAAAGCAAAAGGTAGGTAATGTCAGGGAGTAGTCTTCATGCCTCCTTCA ACTGGGAGCATGTTCTGAGGGTGCCCTCCCAAGCCTGGGAGTAACTATTTCCCCCATCCC CAGGCCTGTGCCCCTCTCTGGTCTCGTGCTTGTGGCAGCTCTGTCTTCAGTTCTGGGATA TGTGCCCGTGTGGATGCTTCATTCCAGCCTCAGGGAAGCCTGGCACCCACTGCCCAACGT GGGCAAAGCGGTATGATGCCTGGCAAAGGGCCTGCATGGCTATCCTCATTGCTACCTAAT GTGCTTGCAAAAGCTCCATGTTTCCTAACAGATTCAGACTCCTGGCCAGGTGTGGTGGCC CACACCTGTAATTCTAGCACTTTGGGAGGCCAAGGTGGGCAGATCACTTGAGGTCAGGAG AAAATTAGCTGGGTGCGCTAGTGCATGCCTGTAATCTCATCTACTCGGGAGGCTAAGACA GGAGACTCTCACTTCAACCCAGGAGGTGGAGGTTGCGGTGAGCCAAGATTGTGCCTCTGC ACTCTAGCGTGGGTGACAGAGTAAGCGAGACTCCATCTCAAAAATAATAATAATAATAAT TCAGACTCCTTATCAGGAGTCCATGATCTGGCCTGGCACAGTAACTCATGCCTGTAATCC CAACATTTTGGGAGGCCAACGCAGGAGGATTGCTTGAGGTCTGGAGGTTTGAGACCAGCC TGGGCAACATAGAAAGACCCCATCTCTAAATAAATGTTTTAAAAAT

FIGURE 126

MELPFVTHLFLPLVFLTGLCSPFNLDEHHPRLFFGPPEAEFGYSVLQHVGGGQRWMLVGA PWDGPSGDRRGDVYRCPVGGAHNAPCAKGHLGDYQLGNSSHPAVNMHLGMSLLETDGDGG FMVS

Important features: Signal peptide: amino acids 1-22

Cell attachment sequence: amino acids 70-73

N-glycosylation site: amino acids 98-101

Integrins alpha chain proteins: amino acids 67-81

FIGURE 127

GAGAGGACGAGGTGCCGCTGCCTGGAGAATCCTCCGCTGCCGTCGGCTCCCGGAGCCCAG CCCTTTCCTAACCCAACCCAACCTAGCCCAGTCCCAGCCGCCAGCGCCTGTCCCTGTCAC GGACCCCAGCGTTACCATGCATCCTGCCGTCTTCCTATCCTTACCCGACCTCAGATGCTC $\overline{\texttt{CCTTCTGCTCCTGGTAACTTGGGTTTTTACTCCTGTAACAACTGAAATAACAAGTCTTGC}$ TACAGAGAATATAGATGAAATTTTAAACAATGCTGATGTTGCTTTAGTAAATTTTTATGC TGACTGGTGTCGTTTCAGTCAGATGTTGCATCCAATTTTTGAGGAAGCTTCCGATGTCAT TAAGGAAGAATTTCCAAATGAAAATCAAGTAGTGTTTGCCAGAGTTGATTGTGATCAGCA CTCTGACATAGCCCAGAGATACAGGATAAGCAAATACCCAACCCTCAAATTGTTTCGTAA TGGGATGATGAAGAGAGAATACAGGGGTCAGCGATCAGTGAAAGCATTGGCAGATTA CATCAGGCAACAAAAAAGTGACCCCATTCAAGAAATTCGGGACTTAGCAGAAATCACCAC TCTTGATCGCAGCAAAAGAAATATCATTGGATATTTTGAGCAAAAGGACTCGGACAACTA TGGGGATGTTTCAAAACCGGAAAGATATAGTGGCGACAACATAATCTACAAACCACCAGG GCATTCTGCTCCGGATATGGTGTACTTGGGAGCTATGACAAATTTTGATGTGACTTACAA TTGGATTCAAGATAAATGTGTTCCTCTTGTCCGAGAAATAACATTTGAAAATGGAGAGGA ATTGACAGAAGAAGGACTGCCTTTTCTCATACTCTTTCACATGAAAGAAGATACAGAAAG TTTAGAAATATTCCAGAATGAAGTAGCTCGGCAATTAATAAGTGAAAAAGGTACAATAAA CTTTTTACATGCCGATTGTGACAAATTTAGACATCCTCTTCTGCACATACAGAAAACTCC CAAAGATGTATTAATTCCTGGAAAACTCAAGCAATTCGTATTTGACTTACATTCTGGAAA ACTGCACAGAGAATTCCATCATGGACCTGACCCAACTGATACAGCCCCAGGAGAGCAAGC CCAAGATGTAGCAAGCAGTCCACCTGAGAGCTCCTTCCAGAAACTAGCACCCAGTGAATA TAGGTATACTCTATTGAGGGATCGAGATGAGCTTTAAAAAACTTGAAAAACAGTTTGTAAG CCTTTCAACAGCAGCATCAACCTACGTGGTGGAAATAGTAAACCTATATTTTCATAATTC ΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑ

FIGURE 128

MHPAVFLSLPDLRCSLLLLVTWVFTPVTTEITSLATENIDEILNNADVALVNFYADWCRF SQMLHPIFEEASDVIKEEFPNENQVVFARVDCDQHSDIAQRXRISKYPTLKLFRNCMMK REYRGQRSVKALADYIRQQKSDPIQEIRDLAEITTLDRSKRNIIGYFEQKDSDNYRVFER VANILHDDCAFLSAFGDVSKPERYSGDNIIYKPPGHSAPDMVYLGAMTNFDVTYNWIQDK CVPLVREITFENGEELTEEGLPFIILFHMKEDTESLEIFQNEVARQLISEKGTINFLHAD CDKFRHPLLHIQKTPADCPVIAIDSFRHMYVFGDFKDVLIPGKLKQFVFDLHSGKLHREF HHGPDPTDTAPGEOAODVASSPPESSFQKLAPSEYRYTLLRDRDEL

Important features: Signal peptide: amino acids 1-29

Endoplasmic reticulum targeting sequence: amino acids 403-406

Tyrosine kinase phosphorylation site: amino acids 203-211

Thioredoxin family proteins: amino acids 50-66

FIGURE 129

GAGCAGGACGGAGCCATGGACCCCGCCAGGAAAGCAGGTGCCCAGGCCATGATCTGGACT CCACCTCCTCCTCCTCCTCCTCCTCCCGCAGGCCCCAGGCCCTGGAGTGCTACAGC TGCGTGCAGAAAGCAGATGACGGATGCTCCCCGAACAAGATGAAGACAGTGAAGTGCGCG CCGGGCGTGGACGTCTGCACCGAGGCCGTGGGGGCGGTGGAGACCATCCACGGACAATTC TCGCTGGCAGTGCGGGGTTGCGGTTCGGGACTCCCCGGCAAGAATGACCGCGGCCTGGAT CTTCACGGGCTTCTGGCGTTCATCCAGCTGCAGCAATGCGCTCAGGATCGCTGCAACGCC AAGCTCAACCTCACCTCGCGGGCGCTCGACCCGGCAGGTAATGAGAGTGCATACCCGCCC AACGGCGTGGAGTGCTACAGCTGTGTGGGCCTGAGCCGGGAGGCGTGCCAGGGTACATCG CCGCCGGTCGTGAGCTGCTACAACGCCAGCGATCATGTCTACAAGGGCTGCTTCGACGGC AACGTCACCTTGACGGCAGCTAATGTGACTGTGTCCTTGCCTGTCCGGGGCTGTGTCCAG GATGAATTCTGCACTCGGGATGGAGTAACAGGCCCAGGGTTCACGCTCAGTGGCTCCTGT TGCCAGGGGTCCCGCTGTAACTCTGACCTCCGCAACAAGACCTACTTCTCCCCTCGAATC CCACCCCTTGTCCGGCTGCCCCCTCCAGAGCCCACGACTGTGGCCTCAACCACATCTGTC ACCACTTCTACCTCGGCCCCAGTGAGACCCACATCCACCAAACCCATGCCAGCGCCA ACCAGTCAGACTCCGAGACAGGGAGTAGAACACGAGGCCTCCCGGGATGAGGAGCCCAGG TTGACTGGAGGCGCCGCTGGCCACCAGGACCGCAGCAATTCAGGGCAGTATCCTGCAAAA GGGGGGCCCCAGCAGCCCCATAATAAAGGCTGTGTGGCTCCCACAGCTGGATTGGCAGCC $\tt CTTCTGTTGGCCGTGGCTGCTGGTGTCCTACTGTGAGCTTCTCCACCTGGAAATTTCCCT$ CTCACCTACTTCTCTGGCCCTGGGTACCCCTCTTCTCATCACTTCCTGTTCCCACCACTG GACTGGGCTGGCCCAGCCCCTGTTTTTCCAACATTCCCCAGTATCCCCAGCTTCTGCTGC GCTGGTTTGCGGCTTTGGGAAATAAAATACCGTTGTATATATTCTGCCAGGGGTGTTCTA GCTTTTTGAGGACAGCTCCTGTATCCTTCTCATCCTTGTCTCTCCGCTTGTCCTCTTGTG TGGCTCCCCACTCTAAGCACTGCCTCCCCTACTCCCCGCATCTTTGGGGAATCGGTTCCC CATATGTCTTCCTTACTAGACTGTGAGCTCCTCGAGGGGGGGCCCGGTACCCAATTCGCC CTATAGTGAGTCGTA

FIGURE 130

MDPARKAGAQAMIWTAGWLLLLLLRGGAQALECYSCVQKADDGCSPNKMKTVKCAPGVDV
CTEAVGAVETIHGGFSLAVRGCCSGLPGKNDRGLDLHGLLAFIQLQQCAQDRCNAKLNLV
SRALDPAGNESAYPPNGVECYSCVGLSREACQGTSPPVASCYNASDHVYKGGFDGNVTLT
AANVTVSLPVRGCVQDEFCTRDGVTGPGFTLSGSCCQGSRCNSDLRNKTYFSPRIPPLVR
LPPPEPTTVASTTSVTTSTSAVVRPTSTTKPMPAPTSQTPRQSVEHEASRDEEPRLTGGA
AGHQDRSNSGQYPAKGGPQQPINKGCVAPTAGLAALLLAVAAGVLL

FIGURE 131

FIGURE 132

MKIPVLPAVVLLSLLVLHSAQGATLGGPEEESTIENYASRPEAFNTPFLNIDKLRSAFKA DEFLNWHALFESIKRKLPFLNWDAFPKLKGLRSATPDAQ

FIGURE 133

FIGURE 134

MGVEIAFASVILTCLSLLAAGVSQVVLLQPVPTQETGPKAMGDLSCGFAGHS

FIGURE 135

GGGGAATCTGCAGTAGGTCTGCCGGCGATGGAGTGGTGGGCTAGCTCGCCGCTTCGGCTC TGGCTGCTGTTGTTCCTCCTGCCCTCAGCGCCAGGGCCCCCAGAAGGAGTCAGGTTCAAAA TGGAAAGTATTTATTGACCAAATTAACAGGTCTTTGGAGAATTACGAACCATGTTCAAGT CAAAACTGCAGCTGCTACCATGGTGTCATAGAAGAGGATCTAACTCCTTTCCGAGGAGGC ATCTCCAGGAAGATGATGGCAGAGGTAGTCAGACGGAAGCTAGGGACCCACTATCAGATC ACTAAGAACAGACTGTACCGGGAAAATGACTGCATGTTCCCCTCAAGGTGTAGTGGTGTT GAGCACTTTATTTTGGAAGTGATCGGGCGTCTCCCTGACATGGAGATGGTGATCAATGTA CGAGATTATCCTCAGGTTCCTAAATGGATGGAGCCTGCCATCCCAGTCTTCTCCTTCAGT AAGACATCAGAGTACCATGATATCATGTATCCTGCTTGGACATTTTGGGAAGGGGGACCT GCTGTTTGGCCAATTTATCCTACAGGTCTTGGACGGTGGGACCTCTTCAGAGAAGATCTG GTAAGGTCAGCAGCACAGTGGCCATGGAAAAAGAAAAACTCTACAGCATATTTCCGAGGA TCAAGGACAAGTCCAGAACGAGATCCTCTCATTCTTCTGTCTCGGAAAAAACCCAAAACTT GTTGATGCAGAATACACCAAAAACCAGGCCTGGAAATCTATGAAAGATACCTTAGGAAAG CCAGCTGCTAAGGATGTCCATCTTGTGGATCACTGCAAATACAAGTATCTGTTTAATTTT CGAGGCGTAGCTGCAAGTTTCCGGTTTAAACACCTCTTCCTGTGTGGCTCACTTGTTTTC CATGTTGGTGATGAGTGGCTAGAATTCTTCTATCCACAGCTGAAGCCATGGGTTCACTAT ATCCCAGTCAAAACAGATCTCTCCAATGTCCAAGAGCTGTTACAATTTGTAAAAGCAAAT GATGATGTAGCTCAAGAGATTGCTGAAAGGGGAAGCCAGTTTATTAGGAACCATTTGCAG ATGGATGACATCACCTGTTACTGGGAGAACCTCTTGAGTGAATACTCTAAATTCCTGTCT TATAATGTAACGAGAAGGAAAGGTTATGATCAAATTATTCCCAAAATGTTGAAAACTGAA CTATAGTAGTCATCATAGGACCATAGTCCTCTTTGTGGCAACAGATCTCAGATATCCTAC GGTGAGAAGCTTACCATAAGCTTGGCTCCTATACCTTGAATATCTGCTATCAAGCCAAAT ACCTGGTTTTCCTTATCATGCTGCACCCAGAGCAACTCTTGAGAAAGATTTAAAATGTGT CTAATACACTGATATGAAGCAGTTCAACTTTTTGGATGAATAAGGACCAGAAATCGTGAG ATGTGGATTTTGAACCCAACTCTACCTTTCATTTTCTTAAGACCAATCACAGCTTGTGCC TCAGATCATCCACCTGTGTGAGTCCATCACTGTGAAATTGACTGTGTCCATGTGATGATG CCCTTTGTCCCATTATTTGGAGCAGAAAATTCGTCATTTGGAAGTAGTACAACTCATTGC TATGGGGTTTATGAAAAATACTTGGGGATCATTCTCTGAATGGTCTAAGGAAGCGGTAGC CATGCCATGCAATGATGTAGGAGTTCTCTTTTGTAAAACCATAAACTCTGTTACTCAGGA GGTTTCTATAATGCCACATAGAAAGAGGCCAATTGCATGAGTAATTATTGCAATTGGATT TCAGGTTCCCTTTTTGTGCCTTCATGCCCTACTTCTTAATGCCTCTCTAAAGCCAAA

FIGURE 136

MEWWASSPLRLWLLLFLLPSAQGRQKESGSKWKVFIDQINRSLENYEPCSSQNCSCYHGV IEEDLTPFRGGISRKMMAEVVRRKLGTHYQITKNRLYRENDCMFPSRCSGVEHFILEVIG RLPDMEMVINVRDYPQVPKWMEPAIPVFSFSKTSEYHDIMYPAWTFWEGGPAVWPIYPTG LGRWDLFREDLVRSAAQWPWKKRNSTAYFRGSRTSPERDPLILLSRKNPKLVDAEYTKNQ AWKSMKDTLGKPAKDVHLVDHCKYKYLFNFRGVAASFRFKHLFLCGSLVFHVGDEWLEF FYPQLKPWVHYIPVKTDLSNVQELLQFVKANDDVAQEIAERGSQFIRNHLQMDDITCYWE NLLSEYSKFLSYNVTRRGYDQIIPKMLKTEL

FIGURE 137

ATTCTCCTAGAGCATCTTTGGAAGCATGAGGCCACGATGCTGCATCTTGGCTCTTGTCTG
CTGGATAACAGTCTTCCTCCTCCAGTGTTCAAAAGGAACTACAGAGGCTCCTGTTGGCTC
AGGACTGTGGCTGTGCCAGCCGACCCAGGTGTGGGAACAAGATCTACAACCCTTCAGA
GCAGTGCTGTTATGATGATGCCATCTTATCCTTAAAGGAGACCCGCCGCTGTGGCTCCAC
CTGCACCTTCTGGCCCTGCTTTGAGCTCTGCTGTCCCGAGTCTTTTGGCCCCCAGCAGAA
GTTTCTTGTGAAGTTGAGGGTTCTGGGTATGAAGTCTCAGTGTCACTTATCTCCCATCTC
CCGGAGCTGTACCAGGAACAGGAGCCACGTCCTGTACCCATAAAAACCCCAGGCTCCACT
GCACACGCCAGACAAGGGAGAAGAGAGACGAACTGACTCAGTTGACCTTACAGTTGA
ACTTCGGAGAAGCAACTTGACTTCAGAGGGATGCCTCAATGACATAGCTTTGGAGAGA
ACCCCAGCTGGGCAGACAAGCCAGCTTCAGGGAAAAGACCCACTTTCCCCTTTC
CAGCTCCCCTTCCCCCTTGAGAGCCACCTTTCATCGGCAATAAAATCCCCCACATTTACCATCT

FIGURE 138

MRPRCCILALVCWITVFLLQCSKGTTDAPVGSGLWLCQPTPRCGNKIYNPSEQCCYDDAI LSLKETRRGGSTCTFWPCFELCCPESFGPQQKFLVKLRVLGMKSQCHLSPISRSCTRNRR HVLVP

Important features: Signal sequence: amino acids 1-21

N-myristoylation sites: amino acids 33-39, 70-76

FIGURE 139

FIGURE 140

MKFTIVFAGLLGVFLAPALANYNINVNDDNNNAGSGQQSVSVNNEHNVANVDNNNGWDSW NSINDYGNGFAATRLFQKKTCIVHKMNKEVMPSIQSLDALVKEKKLQGKGPGGPPFKGLM YSVNPNKVDDLSKFGKNIANMCRGIPTYMAEEMQEASLFFYSGTCYTTSVLWIVDISFCG DTVFN

Signal Peptide: amino acids 1-20

N-myristoylation Sites: amino acids 67-72, 118-123, 163-168

Flavodoxin protein homology: amino acids 156-174

FIGURE 141

GGTCCTTAATGGCAGCAGCCGCCGCTACCAAGATCCTTCTGTGCCTCCCGCTTCTGCTCC TGCTGTCCGGCTGGTCCCGGGCTGGGCGAGCCGACCCTCACTCTTTTGCTATGACATCA CCGTCATCCCTAAGTTCAGACCTGGACCACGGTGGTGTGCGGTTCAAGGCCAGGTGGATG AAAAGACTTTTCTTCACTATGACTGTGGCAACAAGACAGTCACACCTGTCAGTCCCCTGG GGAAGAACTAAATGTCACAACGGCCTGGAAAGCACAGAACCCAGTACTGAGAGAGGTGG TGGACATACTTACAGAGCAACTGCGTGACATTCAGCTGGAGAATTACACACCCAAGGAAC CCCTCACCCTGCAGGCAAGGATGTCTTGTGAGCAGAAAGCTGAAGGACACAGCAGTGGAT CTTGGCAGTTCAGTTTCGATGGGCAGATCTTCCTCCTCTTTGACTCAGAGAAGAGAATGT GGACAACGGTTCATCCTGGAGCCAGAAAGATGAAAGAAAAGTGGGAGAATGACAAGGTTG TGGCCATGTCCTTCCATTACTTCTCAATGGGAGACTGTATAGGATGGCTTGAGGACTTCT TGATGGGCATGGACACCCTGGAGCCAAGTGCAGGAGCACCACTCGCCATGTCCTCAG GCACAACCCAACTCAGGGCCACAGCCACCCTCATCCTTTGCTGCCTCCTCATCATCC TCCCCTGCTTCATCCTCCCTGGCATCTGAGGAGAGTCCTTTAGAGTGACAGGTTAAAGCT TGCCCACGACCTACGGTGTATGTCCAGTGGCCTCCAGCAGATCATGATGACATCATGGAC CCAATAGCTCATTCACTGCCTTGATTCCTTTTGCCAACAATTTTACCAGCAGTTATACCT AACATATTATGCAATTTTCTCTTGGTGCTACCTGATGGAATTCCTGCACTTAAAGTTCTG TCTTTGAATGATGATCTCTTTCTTGCAAATGATATTGTCAGTAAAATAATCACGTTAGAC TTCAGACCTCTGGGGATTCTTTCCGTGTCCTGAAAGAGAATTTTTAAATTATTAATAAG AAAAAATTTATTAATGATTGTTTCCTTTAGTAATTTATTGTTCTGTACTGATATTTAA

FIGURE 142

MAAAAATKILLCLPLLLLLSGWSRAGRADPHSLCYDITVIPKFRPGPRWCAVQGQVDEKT FLHYDCGNKTVTPVSPLGKKLNVTTAWKAQNPVLREVVDILTEQLRDIQLENYTPKEPLT LQARMSCEQKAEGHSSGSWQFSFDGQIFLLFDSEKRWWTTVHPGARKMKEKWENDKVVAM SFHYFSMGDCIGWLEDFLMGMDSTLEPSAGAPLAMSSGTTQLRATATTLILCCLLIILPC FILPGI

Important features: Signal peptide: amino acids 1-25

Transmembrane domain: amino acids 224-246

N-glycosylation site: amino acids 68-72, 82-86

N-myristoylation site: amino acids 200-206, 210-216

Amidation site: amino acids 77-81

FIGURE 143

AATGTGAGAGGGGCTGATGGAAGCTGATAGGCAGGACTGGAGTGTTAGCACCAGTACTGG ATGTGACAGCAGGCAGAGGAGCACTTAGCAGCTTATTCAGTGTCCGATTCTGATTCCGGC AAGGATCCAAGCATGGAATGCTGCCGTCGGGCAACTCCTGGCACACTGCTCCTCTTTCTG GCTTTCCTGCTCCTGAGTTCCAGGACCGCACGCTCCGAGGAGGACCGGGACGGCCTATGG GATGCCTGGGGCCCATGGAGTGAATGCTCACGCACCTGCGGGGGAGGGGCCTCCTACTCT CTGAGGCGCTGCCTGAGCAGCAAGAGCTGTGAAGGAAGAATATCCGATACAGAACATGC AGTAATGTGGACTGCCCACCAGAAGCAGGTGATTTCCGAGCTCAGCAATGCTCAGCTCAT AATGATGTCAAGCACCATGGCCAGTTTTATGAATGGCTTCCTGTGTCTAATGACCCTGAC AACCCATGTTCACTCAAGTGCCAAGCCAAAGGAACAACCCTGGTTGTTGAACTAGCACCT AAGGTCTTAGATGGTACGCGTTGCTATACAGAATCTTTGGATATGTGCATCAGTGGTTTA TGCCAAATTGTTGGCTGCGATCACCAGCTGGGAAGCACCGTCAAGGAAGATAACTGTGGG GTCTGCAACGGAGATGGGTCCACCTGCCGGCTGGTCCGAGGGCAGTATAAATCCCAGCTC TCCGCAACCAAATCGGATGATACTGTGGTTGCACTTCCCTATGGAAGTAGACATATTCGC CTTGTCTTAAAAGGTCCTGATCACTTATATCTGGAAACCAAAACCCTCCAGGGGACTAAA GGTGAAAACAGTCTCAGCTCCACAGGAACTTTCCTTGTGGACAATTCTAGTGTGGACTTC CAGAAATTTCCAGACAAAGAGATACTGAGAATGGCTGGACCACTCACAGCAGATTTCATT GTCAAGATTCGTAACTCGGGCTCCGCTGACAGTACAGTCCAGTTCATCTTCTATCAACCC ATCATCCACCGATGGAGGGAGACGGATTTCTTTCCTTGCTCAGCAACCTGTGGAGGAGGT TATCAGCTGACATCGGCTGAGTGCTACGATCTGAGGAGCAACCGTGTGGTTGCTGACCAA TACTGTCACTATTACCCAGAGAACATCAAACCCAAACCCAAGCTTCAGGAGTGCAACTTG GATCCTTGTCCAGCCAGTGACGGATACAAGCAGATCATGCCTTATGACCTCTACCATCCC CAGAGCCGGGCAGTTTCCTGTGTGGAGGAGGACATCCAGGGGCATGTCACTTCAGTGGAA GAGTGGAAATGCATGTACACCCCTAAGATGCCCATCGCGCAGCCCTGCAACATTTTTGAC TGCCCTAAATGGCTGGCACAGGAGTGGTCTCCGTGCACAGTGACATGTGGCCAGGGCCTC AGATACCGTGTGGTCCTCTGCATCGACCATCGAGGAATGCACACAGGAGGCTGTAGCCCA AAAACAAAGCCCCACATAAAAGAGGAATGCATCGTACCCACTCCCTGCTATAAACCCAAA GAGAAACTTCCAGTCGAGGCCAAGTTGCCATGGTTCAAACAAGCTCAAGAGCTAGAAGAA GGAGCTGCTGTCAGAGGAGCCCTCGTAAGTTGTAAAAGCACAGACTGTTCTATATTTG AAACTGTTTTGTTTAAAGAAAGCAGTGTCTCACTGGTTGTAGCTTTCATGGGTTCTGAAC AAAAAAAA

FIGURE 144

MECCRRATPGTLLLFLAFLLLSSRTARSEEDRDGLWDAWGPWSECSRTCGGGASYSLRRC
LSSKSCEGRNIRYRTCSNVDCPPEAGDFRAQQCSAHNDVKHHGQFYEWLPVSNDPDNPCS
LKCQAKGTTLVVELAPKVLDGTRCYTESLDMCISGLCQIVGCDDQLGSTVKEDDNCGVCNG
DGSTCRLVRGQYKSQLSATKSDDTVVALPYGSRHIRLVLKGPDHLYLETKTLQGTKGENS
LSSTGTFLVDNSSVDFQKFPDKEILRMAGPLTADFIVKIRNSGSADSTVQFIFYQPIIHR
WRETDFFPCSATCGGGYQLTSAECYDLKSNRVVADQVCHYYPENIKPKPLKQECNLDPCP
ASDGYKQIMPYDLYHPLPRWEATPWTACSSSCGGGIGSRAVSCVEEDIQGHVTSVEEWKC
MYTPKMPIAQPCNIFDCPKWLAQEWSPCTVTCGQGLRYRVVLCIDHRGMHTGGCSPKTKP
HIKBECIUPTPCYKPKEKLPVEAKLPWFKQAQELEEGAAVSEEPS

Important features: Signal peptide: amino acids 1-25

N-glycosylation site: amino acids 251-254

Thrombospondin 1: amino acids 385-399

von Willebrand factor type C domain proteins: amino acids 385-399, 445-459 and 42-56

FIGURE 145

GGAGGAGGGAGGCGGCAGGCCCAGCCCAGAGCAGCCCCGGGCACCAGCACGGACTCT CTCTTCCAGCCCAGGTGCCCCCACTCTCGCTCCATTCGGCGGGAGCACCCAGTCCTGTA CGCCAAGGAACTGGTCCTGGGGGCACCATGGTTTCGGCGGCAGCCCCCAGCCTCCTCATC GCCACGTTCCTGGAGGATGTGGCGGGTAGTGGGGAGGCCGAGGGCTCGTCGGCCTCCTCC CCGAGCCTCCCGCCACCCTGGACCCCGGCCCTCAGCCCCACATCGATGGGGCCCCAGCCC CGCCAGTACGTGATGCTGATTGCTGTGGTGGGCTCCCTGGCCTTTCTGCTGATGTTCATC GTCTGTGCCGCGGTCATCACCCGGCAGAAGCAGAAGGCCTCGGCCTATTACCCATCGTCC TTCCCCAAGAAGAAGTACGTGGACCAGAGTGACCGGGCCGGGGCCCCCGGGCCTTCAGT GAGGTCCCCGACAGAGCCCCCGACAGCAGGCCCGAGGAAGCCCTGGATTCCTCCCGGCAG CTCCAGGCCGACATCTTGGCCGCCACCCAGAACCTCAAGTCCCCCACCAGGGCTGCACTG GGCGGTGGGGACGGAGCCAGGATGGTGGAGGGCAGGGGCGCAGAGGAAGAGGAGAAGGGC AGCCAGGAGGGGGACCAGGAAGTCCAGGGACATGGGGTCCCAGTGGAGACACCAGAGGCG CAGGAGGAGCCGTGCTCAGGGGTCCTTGAGGGGGCTGTGGTGGCCGGTGAGGGCCAAGGG GAGCTGGAAGGGTCTCTCTTGTTAGCCCAGGAAGCCCAGGGACCAGTGGGTCCCCCCGAA AGCCCCTGTGCTTGCAGCAGTGTCCACCCCAGTGTCTAACAGTCCTCCCGGGCTGCCAGC CCTGACTGTCGGGCCCCCAAGTGGTCACCTCCCCGTGTATGAAAAGGCCTTCAGCCCTGA CTGCTTCCTGACACTCCCTCCTTGGCCTCCCTGTGGTGCCAATCCCAGCATGTGCTGATT CTACAGCAGGCAGAAATGCTGGTCCCCGGTGCCCCGGAGGAATCTTACCAAGTGCCATCA TCCTTCACCTCAGCAGCCCCAAAGGGCTACATCCTACAGCACAGCTCCCCTGACAAAGTG AGGGAGGCACGTGTCCCTGTGACAGCCAGGATAAAACATCCCCCAAAGTGCTGGGATTA CAGGCGTGAGCCACCGTGCCCGGCCCAAACTACTTTTTAAAACAGCTACAGGGTAAAATC $\tt CTGCAGCACCCACTCTGGAAAATACTGCTCTTAATTTTCCTGAAGGTGGCCCCCTGTTTC$ TAGTTGGTCCAGGATTAGGGATGTGGGGTATAGGGCATTTAAATCCTCTCAAGCGCTCTC CAAGCACCCCGGCCTGGGGGTGAGTTTCTCATCCCGCTACTGCTGCTGGGATCAGGTTG AATGAATGGAACTCTTCCTGTCTGGCCTCCAAAGCAGCCTAGAAGCTGAGGGGCTGTGTT TGAGGGGACCTCCACCCTGGGGAAGTCCGAGGGGCTGGGGAAGGGTTTCTGACGCCCAGC CTGGAGCAGGGGGCCCTGGCCACCCCCTGTTGCTCACACATTGTCTGGCAGCCTGTGTC CACAATATTCGTCAGTCCTCGACAGGGAGCCTGGGCTCCGTCCTGCTTTAGGGAGGCTCT GGCAGGAGGTCCTCTCCCCCATCCCTCCATCTGGGGCTCCCCCAACCTCTGCACAGCTCT

FIGURE 146

MVSAAAPSLLILLLLLLGSVPATDARSVPLKATFLEDVAGSGEAEGSSASSPSLPPPWTP ALSPTSMGPQPTTLGGPSPPTNFLDGIVDFFRQYVMLIAVVGSLAFILMFIVCAAVITRQ KQKASAYYPSSFPKKKYVDQSDRAGGFRAFSEVPDRAPDSRPEEALDSSRQLQADILAAT QNLKSPTRAALGGGDGARMVEGRGAEEEKGSQEGDQEVQGHGVPVETPEAQEEPCSGVL EGAVVAGEGQGELEGSLLLAQEAQGPVGPPESPCACSSVHPSV

Signal peptide: amino acids 1-25

Transmembrane domain: amino acids 94-118

N-myristoylation site: amino acids 18-24, 40-46, 46-52, 145-151, 192-198, 193-199, 211-217, 238-244, 242-248

FIGURE 147

FIGURE 148

MKMLLLLCLGLTLVCVHAEEASSTGRNFNVEKINGEWHTIILASDKREKIEEHGNFRLFL EQIHVLENSLVLKVHTVRDEECSELSMVADKTEKAGEYSVTYDGFNTFTIPKTDYDNFLM AHLINEKDGETFQLMGLYGREPDLSSDIKERFAQLCEEHGILRENIIDLSNANRCLQARE

FIGURE 149

GAGGCTGGCCCCTGCAGAGCGGAGAGGGAAAGCACTGGGACAAATATTGGGGAGGCCCTT GGACATGGCCTGGGAGACGCCCTGAGCGAAGGGGTGGGAAAGGCCATTGGCAAAGAGGCC GGAGGGGCAGCTGGCTCTAAAGTCAGTGAGGCCCTTGGCCAAGGGACCAGAGAAGCAGTT GGCACTGGAGTCAGGCAGGTTCCAGGCTTTGGCGCAGCAGATGCTTTGGGCAACAGGGTC GGGGAAGCAGCCCATGCTCTGGGAAACACTGGGCACGAGATTGGCAGACAGGCAGAAGAT GTCATTCGACACGGAGCAGATGCTGTCCGCGGCTCCTGGCAGGGGGTGCCTGGCCACAGT GGTGCTTGGGAAACTTCTGGAGGCCATGGCATCTTTGGCTCTCAAGGTGGCCTTGGAGGC CAGGGCCAGGGCAATCCTGGAGGTCTGGGGACTCCGTGGGTCCACGGATACCCCGGAAAC TCAGCAGGCAGCTTTGGAATGAATCCTCAGGGAGCTCCCTGGGGTCAAGGAGGCAATGGA GGGCCACCAAACTTTGGGACCAACACTCAGGGAGCTGTGGCCCAGCCTGGCTATGGTTCA GTGAGAGCCAGCAACCAGAATGAAGGGTGCACGAATCCCCCACCATCTGGCTCAGGTGGA GGCTCCAGCAACTCTGGGGGAGGCAGCGGCTCACAGTCGGGCAGCAGTGGCAGTGGCAGC AATGGTGACAACAACAATGGCAGCAGCAGTGGTGGCAGCAGCAGTGGCAGCAGTGGC AGCAGCAGTGGCGGCAGCAGTGGCGGCAGCAGTGGCAGCAGTGGCAACAGTGGTGGC AGCAGAGGTGACAGCGGCAGTGAGTCCTCCTGGGGATCCAGCACCGGCTCCTCCTCCGGC AACCACGGTGGGAGCGGCGGAGGAAATGGACATAAACCCGGGTGTGAAAAGCCAGGGAAT GAAGCCCGCGGGAGCGGGAATCTGGGATTCAGGGCTTCAGAGGACAGGGAGTTTCCAGC AACATGAGGGAAATAAGCAAAGAGGGCAATCGCCTCCTTGGAGGCTCTGGAGACAATTAT CGGGGCAAGGGTCGAGCTGGGGCAGTGGAGGAGGTGACGCTGTTGGTGGAGTCAATACT GTGAACTCTGAGACGTCTCCTGGGATGTTTAACTTTGACACTTTCTGGAAGAATTTTAAA TCCAAGCTGGGTTTCATCAACTGGGATGCCATAAACAAGGACCAGAGAAGCTCTCGCATC AACACCACCCTCTCATCACTAATCTCAGCCCTTGCCCTTGAAATAAACCTTAGCTGCCCC

FIGURE 150

```
Signal peptide:
amino acids 1-21
N-glycosylation site:
amino acids 265-269
```

Glycosaminoglycan attachment site: amino acids 235-239, 237-241, 244-248, 255-259, 324-328, 388-392

Casein kinase II phosphorylation site: amino acids 26-30, 109-113, 259-263, 300-304, 304-308

```
N-myristoylation site:
```

amino acids 17-23, 32-38, 42-48, 50-56, 60-66, 61-67, 64-70, 74-80, 90-96, 96-102, 130-136, 140-146, 149-155, 152-158, 155-161, 159-165, 163-169, 178-184, 190-196, 194-200, 199-205, 218-224, 236-242, 238-244, 239-245, 240-246, 245-251, 246-252, 249-252, 253-259, 256-262, 266-272, 270-276, 271-277, 275-281, 279-285, 283-289, 284-290, 287-293, 288-294, 291-297, 292-298, 295-301, 298-304, 305-311, 311-317, 315-321, 319-325, 322-328, 323-329, 325-331, 343-349, 354-360, 356-362, 374-380, 381-387, 383-389, 387-393, 389-395, 395-401

Cell attachment sequence: amino acids 301-304

FIGURE 151

CGGCCACAGCTGGCATGCTCTGCCTGATCGCCATCCTGCTGTATGTCCTCGTCCAGTACC TCGTGAACCCCGGGGTGCTCCGCACGGACCCCAGATGTCAAGAATATGAACACGTGGCTG CTGTTCCTCCCCTGTTCCCGGTGCAGGTGCAGACCCTGATAGTCGTGATCATCGGGATG CTCGTGCTCCTGCTGGACTTTCTTGGCTTGGTGCACCTGGGCCAGCTGCTCATCTTCCAC ATCTACCTGAGTATGTCCCCCACCCTAAGCCCCCGATCCCCCAAGGCTGGGTGGTCAGA GCTGCTCATCTTACACCTCTACTTGAGTATGTCCCTAACCCTGAGCCCCCCACGCCTGGG GCCAGAGTCTTTGTCCCCCGTGTGCGCATGTGTTCAGGGTCAGCCTCTCCCAGAAGTGAG ATCATGGACAAAAAGGGCAAATCACAGGAAGAAATTAAATCCATGAGGACCCAGCAGGCC CAGCAAGAAGCTGAACTCACGCCGAGACCTGCAGGAGTGGTGCCAGGTGCT<u>TGA</u>AGTAAC AAGTTTAAAATGTTCAGAGACAATGGAATGGAATCTATTAGGCAAGAACAGGACATTATG AAATAAGGACAGGTGGACTTCCAAAAACACAAGTAGAAATTCTAACAATGAAATATATTA CAGTGGGCACAGCGGTAGGCGGTCAGTCATGTTGCTGAACGACGGAGGGTAAACTCCCCA GCCCCAAGAAAACCTGTGTTGGAAGTAACAACAACCTCCCTGCTCCTGGCACCAGCCGTT TTGGTCATGGTGGGCCAGCTGCAAAGCGTCTTCCATTCTCTGGGCAGTGGTGGCCCCGAG GCTGTGGCCTCTCAGGGGGTTTCTGTGGACACGGGCAGCAGAGTGTGTCCAGGCCAGCCC CCAAGAATGCCCTGCTCCTGACAGCTTGGCCAACCCCTGGTCAGGGCAGAGGGAGTTGGG GAACGCCCAGCTCAGAATGAACACACCCCACCAAGAGCCTCCTTGTTCATAACCACAGGT AAAAAAAAAAA

FIGURE 152

MNTWLLFLPLFPVQVQTLIVVIIGMLVLLLDFLGLVHLGQLLIFHIYLSMSPTLSPRSPQ GWVVRAAHLTPLLEYVPMPEPPTPGARVFVPRVRMCSGSASPRSEIMDKKGKSQEEIKSM RTQQAQQEAELTPRPAGVVPGA

FIGURE 153

AACTGGAAGGAAAGAAAGAAAGGTCAGCTTTGGCCCAGATGTGGTTACCCCTTGGTCTCC $\overline{\text{TGTCTTTATGTCTTTCTCCTCTTTCCTATTCTGTCATCTCCCTCACTTAAGTCTCAGGCCT}$ GTCAGCAGCTCCTGTGGACATTGCCATCCCCTCTGGTAGCCTTCAGAGCAAACAGGACAA CCTATGTTATGGATGTTTCCACCAACCAGGGTAGTGGCATGGAGCACCGTAACCATCTGT GCTTCTGTGATCTCTATGACAGAGCCACTTCTCCACCTCTGAAATGTTCCCTGCTCTGAA TCCTCCTCCCAAGTCTGTTCTCTTATTGTCAACCTCAGCACAACAGGCTGGCGCCAATGG CATTACAGAGAAAGCAATCTGTGTGGCTAGTGGGCAGATTACCATGCAAGCCCCAGGAGA AATGGAGGAGCTTTGTAGCCACCTCCCTGTCAGCCAGTATTAACATGTCCCCTTCCCCCT GCCCCGCCGTAGATTCAGGACATTCGCCCCTGTGTGCCACCAAACCAGGACTTTCCCCTT GGCTTGGCATCCCTGGCTCTCCTGGTACCCAGCAAGACGTCTGTTCCAGGGCAGTGTA GCATCTTTCAAGCTCCGTTACTATGGCGATGGCCATGATGTTACAATCCCACTTGCCTGA ATAATCAAGTGGGAAGGGGAAGCAGAGGGAAATGGGGCCATGTGAATGCAGCTGCTCTGT TCTCCCTACCCTGAGGAAAAACCAAAGGGAAGCAACAGGAACTTCTGCAACTGGTTTTTA TCGGAAAGATCATCCTGCCTGCAGATGCTGTTGAAGGGGCACAAGAAATGTAGCTGGAGA AGATTGATGAAAGTGCAGGTGTGTAAGGAAATAGAACAGTCTGCTGGGAGTCAGACCTGG AATTCTGATTCCAAACTCTTTATTACTTTGGGAAGTCACTCAGCCTCCCCGTAGCCATCT CCAGGGTGACGGAACCCAGTGTATTACCTGCTGGAACCAAGGAAACTAACAATGTAGGTT ACTAGTGAATACCCCAATGGTTTCTCCAATTATGCCCATGCCACCAAAACAATAAAACAA AATTCTCTAACACTGAAA

FIGURE 154

MWLPLGLLSLCLSPLPILSSPSLKSQACQQLLWTLPSPLVAFRANRTTYVMDVSTNQGSG MEHRNHLCFCDLYDRATSPPLKCSLL

FIGURE 155

CAGGAGCGACGTCACCGCC**ATG**GCAGGCATCAAAGCTTTGATTAGTTTGTCCTTTGGAGG AGCAATCGGACTGATGTTTTTGATGCTTGGATGTGCCCTTCCAATATACAACAAATACTG GCCCCTCTTTGTTCTATTTTTTTACATCCTTTCACCTATTCCATACTGCATAGCAAGAAG AACGGGCATTGTCGTGTCAGCTTTTGGACTCCCTATTGTATTTGCCAGAGCACATCTGAT TGAGTGGGGAGCTTGTGCACTTGTTCTCACAGGAAACACAGTCATCTTTGCAACTATACT AGGCTTTTTCTTGGTCTTTGGAAGCAATGACGACTTCAGCTGGCAGCAGTGGTGAAAAAGA AATTACTGAACTATTGTCAAATGGACTTCCTGTCATTTGTTGGCCATTCACGCACACAGG AGATGGGGCAGTTAATGCTGAATGGTATAGCAAGCCTCTTGGGGGTATTTTAGGTGCTCC CTTCTCACTTTTATTGTAAGCATACTATTTTCACAGAGACTTGCTGAAGGATTAAAAGGA TTTTCTCTTTTGGAAAAGCTTGACTGATTTCACACTTATCTATAGTATGCTTTTTGTGGT ATATGCAATGTTAAACACTTTTTTAATGTAATCATTTGCATTGGTTAGGAATTCAGAATT CCGCCGGCTCTATTACTGGTCAAGTACATCTTTTCTCTTAAAATTATTTAGCCTCCATTA TTACAAAAAATTATAAAAATAAGTTTTCAGTCAGTCAGGATGACATCACTCCCAATGTTA TGCAGACATACAGACGGTTGGCATACGTTATAGACTGTATACTCAGTGCAAATATAGCTG CATTTATACCTCAGAGGGGCCAAGTGTTAATGCCCATGCCCTCCGTTAAGGGTTGTTGGT TTTACTGGTAGACAGATGTTTTGTGGATTGAAAATTATTTTATGGAATTGCTACAGAGGA GTGCTTTTCTTCTCAATTGTTAGAAGAATTTATGTTAAACTTTAAGGTAAGGGTGTAAAA ACATTTTTGAGATAAGGTTTTTATTTATGTTTATTATTGTTAGAGTGAGTTGCAATGTGG GAAGAAATGACATTGAAATTCCAGTTTTTGAATCCTGTTTCTATTTATAAGTGAAATTTG CTGATGAGGGACAGTTGTATGTTTGCATCATATATGCCAGAAAACCTTCCTCTGCTTCCT CCTTTTGACTTATTTGGTATGTTGTATATATTACATAAAATAACTTTTCAAATATAGTTT AATAACACTTAGAAGTGTTTACTTACCTGGAAAATAATTGCTATGCCGTACATTCAGAGT GCCCCTCCCCTGCAAGGCCTTGCCATGATTAACAAGTAACTTGTTAGTCTTACAGATAA TTCATGCATTAACAGTTTAAGATTTAGACCATGGTAATAGTAGTTCTTATTCTCTAAGGT TATATCATATGTAATTTAAAAGTATTTTTAAGACAAGTTTCCTGTATACCTCTGAACTGT TTTGATTTTGAGTTCATCATGATAGATCTGCTGTTTCCTTATAAAAGGCATTTGTTGTGT GAGTTAATGCAAAGTAGCCAAGTCCAGCTATATAGCAGCTTCAGAAACATACCTGACCAA AAAATTCCCAGTAACCAGGCATGATCAATTTATAGTGGTCGTTTACATCTAATAATTATC AGGACTTTTTTCAGGAGTGGGTTATAAAAACATTCAAGTTGGTCTGACAGTATTTTGTTA AGGATATTTGTTTGTATGTTTATTCAGTATACTTACATAAAAATTATTTCGCCATCAGCC AAAACTCAGTAATCATGACAGCTGTCTGTTGTTTTATGAAGTTTATTTCTCAAGAAAATG GGAATAAATTTGGGATTTGTTCAGCTTTTTTACTAAAGATGCCTAAAGCCACAGGTTTTA TTGCCTAACTTAAGCCATGACTTTTAGATATGAGATGACGGGAAGCAGGACGAAATATCG GCGTGTGGCTGGAGCCTTCCCACTGGAGGCTGAAAGTGGCTTGTGGTATTATAATGTTCA GATTTCAAGAGGAAGGTGCAGGTACACATGAGTTAGAGAGCTGGTGAGACAGTTGGGAAC TCTTTGTGCTTGTGATCTACTGGACTTTTTTTTTGCAGGAAGTGCATTCTCTGGTCCTTC CCTATTTTCTGTTCTGGATGTCAGTGCAGTGCACTGCTACTGTTTTATCCACTTGGCCAC AGACTTTTTCTAACAGCTGCGTATTATTTCTATATACTAATTGCATTGGCAGCATTGTGT CTTTGACCTTGTATACTAGCTTGACATAGTGCTGTCTCTGATTTCTAGGCTAGTTACTTG AGATATGAATTTTCCATAGAATATGCACTGATACAACATTACCATTCTTCTATGGAAAGA

FIGURE 156

MAGIKALISLSFGGAIGLMFLMLGCALPIYNKYWPLFVLFFYILSPIPYCIARRLVDDTD AMSNACKELAIFLTTGIVVSAFGLPIVFARAHLIEWGACALVLTGNTVIFATILGFFLVF GSNDDFSWOOW

FIGURE 157

GTTTCTCATAGTTGGCGTCTTCTAAAGGAAAAACACTAAAATGAGGAACTCAGCGGACCG GGAGCGACGCAGCTTGAGGGAAGCATCCCTAGCTGTTGGCGCAGAGGGGCGAGGCTGAAG CCGAGTGGCCCGAGGTGTCTGAGGGGCTGGGGCAAAGGTGAAAGAGTTTCAGAACAAGCT TCCTGGAACCCATGACCCATGAAGTCTTGTCGACATTTATACCGTCTGAGGGTAGCAGCT CGAAACTAGAAGAAGTGGAGTGTTGCCAGGGACGGCAGTATCTCTTTGTGTGACCCTGGC GGCCTATGGGACGTTGGCTTCAGACCTTTGTGATACACCATGCTGCGTGGGACGATGACG GCGTGGAGAGGAATGAGGCCTGAGGTCACACTGGCTTGCCTCCTCCTAGCCACAGCAGGC TGCTTTGCTGACTTGAACGAGGTCCCTCAGGTCACCGTCCAGCCTGCGTCCACCGTCCAG AAGCCCGGAGGCACTGTGATCTTGGGCTGCGTGGTGGAACCTCCAAGGATGAATGTAACC TGGCGCCTGAATGGAAAGGAGCTGAATGGCTCGGATGATGCTCTGGGTGTCCTCATCACC CACGGGACCCTCGTCATCACTGCCCTTAACAACCACACTGTGGGACGGTACCAGTGTGTG CAGGACTTCAAGTTAGATGTGCAGCACGTGATTGAAGTGGATGAGGGAAACACAGCAGTC ATTGCCTGCCACCTGCTGAGAGCCACCCCAAAGCCCAGGTCCGGTACAGCGTCAAACAA GAGTGGCTGGAGGCCTCCAGAGGTAACTACCTGATCATGCCCTCAGGGAACCTCCAGATT GTGAATGCCAGCCAGGAGGACGAGGGCATGTACAAGTGTGCAGCCTACAACCCAGTGACC CAGGAAGTGAAAACCTCCGGCTCCAGCGACAGGCTACGTGTGCGCCGCTCCACCGCTGAG GCTGCCCGCATCATCTACCCCCCAGAGGCCCAAACCATCATCGTCACCAAAGGCCAGAGT CTCATTCTGGAGTGTGGGCCAGTGGAATCCCACCCCCACGGGTCACCTGGGCCAAGGAT GGGTCCAGTGTCACCGGCTACAACAAGACGCGCTTCCTGCTGAGCAACCTCCTCATCGAC ACCACCAGCGAGGAGGACTCAGGCACCTACCGCTGCATGGCCGACAATGGGGTTGGGCAG CCCGGGGCAGCGTCATCCTCTACAATGTCCAGGTGTTTGAACCCCCTGAGGTCACCATG GAGCTATCCCAGCTGGTCATCCCCTGGGGCCAGAGTGCCAAGCTTACCTGTGAGGTGCGT CGCCTCCGGCTCTCCCGCAGGCCCTGCGCGTGCTCAGCATGGGGCCTGAGGACGAAGGC GTCTACCAGTGCATGGCCGAGAACGAGGTTGGGAGCGCCCATGCCGTAGTCCAGCTGCGG ACCTCCAGGCCAAGCATAACCCCAAGGCTATGGCAGGATGCTGAGCTGGCTACTGGCACA CCTCCTGTATCACCCTCCAAACTCGGCAACCCTGAGCAGATGCTGAGGGGGCAACCGGCG CTCCCCAGACCCCCAACGTCAGTGGGGCCTGCTTCCCCGAAGTGTCCAGGAGAGAGGGG TCATATGAACTGGTGTGGCGGCCTCGGCATGAGGCCAGTGGCCGGGCGCCAATCCTCTAC TATGTGGTGAAACACCGCAAGCAGGTCACAAATTCCTCTGACGATTGGACCATCTCTGGC ATTCCAGCCAACCAGCACCGCCTGACCCTCACCAGACTTGACCCCGGGAGCTTGTATGAA GTGGAGATGGCAGCTTACAACTGTGCGGGAGAGGGCCAGACAGCCATGGTCACCTTCCGA ACTGGACGCCGCCAAACCCGAGATCATGGCCAGCAAAGAGCAGCAGATCCAGAGAGAC GACCCTGGAGCCAGTCCCCAGAGCAGCCAGCCAGACCACGGCCGCCTCTCCCCCCA GAAGCTCCCGACAGGCCCACCATCTCCACGGCCTCCGAGACCTCAGTGTACGTGACCTGG ATTCCCCGTGGGAATGGTGGGTTCCCAATCCAGTCCTTCCGTGTGGAGTACAAGAAGCTA AAGAAAGTGGGAGACTGGATTCTGGCCACCAGCGCCATCCCCCATCGCGGCTGTCCGTG GAGATCACGGGCCTAGAGAAAGGCACCTCCTACAAGTTTCGAGTCCGGGCTCTGAACATG CTGGGGGAGAGCGAGCCCAGCGCCCCTCTCGGCCCTACGTGGTGTCGGGCTACAGCGGT CGCGTGTACGAGAGGCCCGTGGCAGGTCCTTATATCACCTTCACGGATGCGGTCAATGAG ACCACCATCATGCTCAAGTGGATGTACATCCCAGCAAGTAACAACAACACCCCAATCCAT GGCTTTTATATCTATTATCGACCCACAGACAGTGACAATGATAGTGACTACAAGAAGGAT ATGGTGGAAGGGGACAAGTACTGGCACTCCATCAGCCACCTGCAGCCAGAGACCTCCTAC GACATTAAGATGCAGTGCTTCAATGAAGGAGGGGAGAGCGAGTTCAGCAACGTGATGATC TGTGAGACCAAAGCTCGGAAGTCTTCTGGCCAGCCTGGTCGACTGCCACCCCCAACTCTG GCCCACCACAGCCGCCCTTCCTGAAACCATAGAGCGGCCGGTGGGCACTGGGGCCATG GTGGCTCGCTCCAGCGACCTGCCCTATCTGATTGTCGGGGTCGTCCTGGGCTCCATCGTT CTCATCATCGTCACCTTCATCCCCTTCTGCTTGTGGAGGGCCTGGTCTAAGCAAAAACAT ACAACAGACCTGGGTTTTCCTCGAAGTGCCCTTCCACCCTCCTGCCCGTATACTATGGTG CCATTGGGAGGACTCCCAGGCCACCAGGCCAGTGGACAGCCCTACCTCAGTGGCATCAGT

GGACGGCCTGTGCTAATGGGATCCACATGAATAGGGGCTGCCCTCGGCTGCAGTGGGC TACCCGGCATGAAGCCCCAGCAGCACTGCCCAGGCGAGCTTCAGCAGCAGAGTGACACC AGCAGCCTGCTGAGGCAGACCCATCTTGGCAATGGATATGACCCCCAAAGTCACCAGATC ACGAGGGTCCCAAGTCTAGCCCGGACGACGTCTTTCTTATACACACTGCCCGACGAC TCCACTCACCAGCTGCTGCAGCCCCATCACGACTGCTGCCAACGCCAGGAGCAGCCTGCT GCTGTGGGCCAGTCAGGGGTGAGGAGAGCCCCCGACAGTCCTGTCCTGGAAGCAGTGTGG GACCCTCCATTCACTCAGGGCCCCCATGCTGCTTGGGCCTTGTGCCAGTTGAAGAGGTG GACAGTCCTGACTCCTGCCAAGTGAGTGGAGGAGACTGGTGTCCCCAGCACCCCGTAGGG GCCTACGTAGGACAGGAACCTGGAATGCAGCTCTCCCCGGGGCCACTGGTGCGTGTGTCT TTTGAAACACCACCTCTCACAATT<u>TAG</u>GCAGAAGCTGATATCCCAGAAAGACTATATATT TGGAGAGACATAAGGAGTCCTACCCGTTGAGGTTGGAGAGGGAAAATAAAGAAGCTGCCA CCTAACAGGAGTCACCCAGGAAAGCACCGCACAGGCTGGCGGGGACAGACTCCTAACCT GGGGCCTCTGCAGTGGCAGGCGAGGCTGCAGGAGGCCCACAGATAAGCTGGCAAGAGGAA GGATCCCAGGCACATGGTTCATCACGAGCATGAGGGGAACAGCAAGGGGCACGGTATCACA GCCTGGAGACACCCACACAGATGGCTGGATCCGGTGCTACGGGAAACATTTTCCTAAGAT GCCCATGAGAACAGACCAAGATGTGTACAGCACTATGAGCATTAAAAAACCTTCCAGAAT TAGTCTTCCCTGTAAAA

FIGURE 158

MLRGTMTAWRGMRPEVTLACLLLATAGCFADLNEVPOVTVOPASTVOKPGGTVILGCVVE PPRMNVTWRLNGKELNGSDDALGVLITHGTLVITALNNHTVGRYOCVARMPAGAVASVPA TVTLANLODFKLDVOHVIEVDEGNTAVIACHLPESHPKAOVRYSVKOEWLEASRGNYLIM PSGNLOIVNASOEDEGMYKCAAYNPVTQEVKTSGSSDRLRVRRSTAEAARIIYPPEAOTI IVTKGQSLILECVASGIPPPRVTWAKDGSSVTGYNKTRFLLSNLLIDTTSEEDSGTYRCM ADNGVGQPGAAVILYNVOVFEPPEVTMELSOLVIPWGOSAKLTCEVRGNPPPSVLWLRNA VPLISSORLRLSRRALRVLSMGPEDEGVYOCMAENEVGSAHAVVOI.RTSRPSITPRI.WOD AELATGTPPVSPSKLGNPEOMLRGOPALPRPPTSVGPASPKCPGEKGOGAPAFAPITI.SS PRTSKTDSYELVWRPRHEGSGRAPILYYVVKHRKOVTNSSDDWTISGIPANOHRLTLTRL DPGSLYEVEMAAYNCAGEGOTAMVTFRTGRRPKPEIMASKEOOTORDDPGASPOSSSOPD HGRLSPPEAPDRPTISTASETSVYVTWIPRGNGGFPIOSFRVEYKKLKKVGDWILATSAT PPSRLSVEITGLEKGTSYKFRVRALNMLGESEPSAPSRPYVVSGYSGRVYERPVAGPYIT FTDAVNETTIMLKWMYI PASNNNTPIHGFYI YYR PTDSDNDSDYKKDMVEGDKYWHSI SH LOPETSYDIKMOCFNEGGESEFSNVMICETKARKSSGOPGRLPPPTLAPPOPPLPETIER PVGTGAMVARSSDLPYLIVGVVLGSIVLIIVTFIPFCLWRAWSKOKHTTDLGFPRSALPP SCPYTMVPLGGLPGHQASGQPYLSGISGRACANGIHMNRGCPSAAVGYPGMKPQOHCPGE LQQQSDTSSLLRQTHLGNGYDPOSHOITRGPKSSPDEGSFLYTLPDDSTHOLLOPHHDCC QRQEOPAAVGOSGVRRAPDSPVLEAVWDPPFHSGPPCCLGLVPVEEVDSPDSCOVSGGDW CPOHPVGAYVGOEPGMOLSPGPLVRVSFETPPLTI

Signal peptide: amino acids 1-30

Transmembrane domain: amino acids 16-30 (type II), 854-879

FIGURE 159

CCCACGCGTCCGCCCACGCGTCCGCCCACGCGTCCGCCCACGCGTCCGCCCACGCGTCCG CCCACGCGTCCGCCCACGCGTCCGGTGCAAGCTCGCGCCGCACACTGCCTGGTGGAGGGA CGCCCGCCGCCGCCCGCAAAGCATGAGTGAGCCCGCTCTCTGCAGCTGCCCGGGGC GCGAATGGCAGGCTGTTTCCGCGGAGTAAAAGGTGGCGCCGGTCAGTGGTCGTTTCCAAT GACGGACATTAACCAGACTGTCAGATCCTGGGGAGTCGCGAGCCCCGAGTTTGGAGTTTT GCTCGGGCTCCGGCACGTAGTTGGGAAACTTGCGGGTCCTAGAAGTCGCCTCCCCGCCTT GCCGGCCGCCTTGCAGCCCGAGCCGAGCAGCAAAGTGAGACATTGTGCGCCTGCCAGA TCCGCCGGCCGCGGACCGGGGCTGCCTCGGAAACACAGAGGGGTCTTCTCTCGCCCTGCA TATAATTAGCCTGCACACAAAGGGAGCAGCTGAATGGAGGTTGTCACTCTCTGGAAAAGG ATTTCTGACCGAGCGCTTCCAATGGACATTCTCCAGTCTCTCTGGAAAGATTCTCGCTAA TGGATTTCCTGCTGCTCGGTCTCTGTCTATACTGGCTGCTGAGGAGGCCCTCGGGGGTGG AGCTGTGCCGGTGCGAGGGGCGGCTGCTGTACTGCGAGGCGCTCAACCTCACCGAGGCGC CCCACAACCTGTCCGGCCTGCTGGGCTTGTCCCTGCGCTACAACAGCCTCTCGGAGCTGC GCGCCGGCCAGTTCACGGGGTTAATGCAGCTCACGTGGCTCTATCTGGATCACAATCACA TCTGCTCCGTGCAGGGGGACGCCTTTCAGAAACTGCGCCGAGTTAAGGAACTCACGCTGA GTTCCAACCAGATCACCCAACTGCCCAACACCCCTTCCGGCCCATGCCCAACCTGCGCA GCGTGGACCTCTCGTACAACAAGCTGCAGGCGCTCGCGCCCGACCTCTTCCACGGGCTGC GGAAGCTCACCACGCTGCATATGCGGGCCAACGCCATCCAGTTTGTGCCCGTGCGCATCT TCCAGGACTGCCGCAGCCTCAAGTTTCTCGACATCGGATACAATCAGCTCAAGAGTCTGG CGCGCAACTCTTTCGCCGGCTTGTTTAAGCTCACCGAGCTGCACCTCGAGCACAACGACT GGAGGAACAAGGTGGCCATTGTGGTCAGCTCGCTGGACTGGGTTTGGAACCTGGAGAAAA TGGACTTGTCGGGCAACGAGATCGAGTACATGGAGCCCCATGTGTTCGAGACCGTGCCGC ACCTGCAGTCCCTGCAGCTGGACTCCAACCGCCTCACCTACATCGAGCCCCGGATCCTCA ACTCTTGGAAGTCCCTGACAAGCATCACCCTGGCCGGGAACCTGTGGGATTGCGGGCGCA ACGTGTGTGCCCTAGCCTCGTGGCTCAGCAACTTCCAGGGGCGCTACGATGGCAACTTGC AGTGCGCCAGCCCGGAGTACGCACAGGGCGAGGACGTCCTGGACGCCGTGTACGCCTTCC ACCTGTGCGAGGATGGGGCCGAGCCCACCAGCGGCCACCTGCTCTCGGCCGTCACCAACC GCAGTGATCTGGGGCCCCCTGCCAGCTCGGCCACCACGCTCGCGGACGGCGGGGAGGGGC AGCACGACGGCACATTCGAGCCTGCCACCGTGGCTCTTCCAGGCGGCGAGCACGCCGAGA GACAGTGCTTTGTCACGCAGCGCAGGAAGCAAAAGCAGAAACAGACCATGCATCAGATGG CTGCCATGTCTGCCCAGGAATACTACGTTGATTACAAACCGAACCACATTGAGGGAGCCC TGGTGATCATCAACGAGTATGGCTCGTGTACCTGCCACCAGCAGCCCGCGAGGGAATGCG AGGTGTGATTGTCCCAGTGGCTCTCAACCCATGCGCTACCAAATACGCCTGGGCAGCCGG GACGGGCCGGCGGCACCAGGCTGGGGTCTCCTTGTCTGTGCTCTGATATGCTCCTTGAC TGAAACTTTAAGGGGATCTCTCCCAGAGACTTGACATTTTAGCTTTATTGTGTCTTAAAA TTAAATTTCATATGAGAACTCCTTCCTCCCTTTGAAGATCTGTCCATATTCAGGAATCTG AGAGTGTAAAAAAGGTGGCCATAAGACAGAGAGAGAATAATCGTGCTTTGTTTTATGCTA CTCCTCCCACCCTGCCCATGATTAAACATCATGTATGTAGAAGATCTTAAGTCCATACGC ATTTCATGAAGAACCATTGGAAAGAGGAATCTGCAATCTGGGAGCTTAAGAGCAAATGAT

GACCATAGAAAGCTATGTTCTTACTTTGTGTGTGTGTCTGTATGTTTCTGCGTTGTGTGT CTTTGTAGGCAAGCAAACGTTGTCTACACAAACGGGAATTTAGCTCACATCATTTCATGC CCCTGTGCCTCTAGCTCTGGAGATTGGTGGGGGGAGGTGGGGGGAAACGGCAGGAATAAG GGAAAGTGGTAGTTTTAACTAAGGTTTTGTAACACTTGAAATCTTTTCTTCTCAAATTA ATTATCTTTAAGCTTCAAGAAACTTGCTCTGACCCCTCTAAGCAACTACTAAGCATTTA CTAATCTCATTATGCTGTGCTATCTGAAAAGAACTTAAGGCCACAATTCACGTCTCGTCC TGGGCATTGTGATGGATTGACCCTCCATTTGCAGTACCTTCCCAGCTGATTAAAGTTCAG CAGTGGTATTGAGGTTTTTCGAATATTTATATAGAAAAAAGTCTTTTCACATGACAAAT GACACTCTCACACCAGTCTTAGCCCTAGTAGTTTTTTAGGTTGGACCAGAGGAAGCAGGT TAAATGAGACCTGTCCTCTGCTGCACTCAGAAAAAATAGGCAGTCCCTGATGCTCAGATC TTAGCCTTGATATTAATAGTTGAGACCACCTACCCACAATGCAGCCTATACTCCCAAGAC TACAAAGTTACCATCGCAAAGGAAAGGTTATTCCAGTAAAAGGAAATAGTTTTCTCAACC ATTTAAAAATATTCTTCTGAACTCATCAAAGTAGAAGAGCCCCCAACCTTTTCTCTCTGC CTTCAAGAAGGCAGACATTTGGTATGATTTAGCATCAACAACACATTTATGAGTATATGT ACAGATCTCTGGTAGGATTAGGGGCCACTTGTGTTTCCGGCTTATTTTAGTCGACTTGTC AGCAAGTTTGATGCCTAGTCTATCTGACATGGCCCAGTAGAACAGGGCATTGATGGATCA CATGAGATGGTAGAAGGAACATCATCACATACCCCTCTCACAGAGAAAATTATCAAAGAA CCAGAAATTATATCTGTTTTGGAGCAAGAGTGTCATAATGTTTCAGGGTAGTCAAAATAA ACATAAATTATCTCCTCTAGATGAGTGGCGATGTTGGCTGATTTGGGTCTGCCATTGACA GAATGTCAAATAAAAAGGAATTAGCTAGAATATGACCATTAAATGTGCTTCTGAAATATA TTTTCACATACCTTTAGAATGTCA

FIGURE 160

MDFLLLGLCLYWLLRRPSGVVLCLLGACFQMLPAAPSGCPQLCRCEGRLLYCEALNLTEA
PHNLSGLLGLSLRYNSLSELRAGQFTGLMQLTWLYYLDHNHICSVQGDAFQKLRRVEELTL
SSNQITQLPNTTFRPMPNLRSVDLSYNKLQALAPPLFHGLRKLTTLHMRANAIOFVPVRI
FQDCRSLKFLDIGYNQLKSLARNSFAGLFKLTELHLEHNDLVKVNFAHFPRLISLHSLCL
RRNKVAIVVSSLDWVWNLEKMDLSGNEIEYMEPHVFETVPHLQSLQLDSNRLTYIEPRIL
NSWKSLTSITLAGRUMDCGRNVCALASMLSNFGGRYPGNLQCASPEYAQGEDVLDAVYAF
HLCEDGAEPTSGHLLSAVTNRSDLGPPASSATTLADGGEGQHOGTFEPATVALPGGEHAE
NAVQIHKVVTGTMALIFSFLIVVLVLVYSWKCFPASLRQLRQCFVTQRRKQKQKQTMHQM
AAMSAOEYYDYKPNHIEGALVIINEYGSCTCHQQPARECEV

FIGURE 161

GGCCGCCTGGAATTGTGGGAGTTGTGTCTGCCACTCGGCTGCCGGAGGCCGAAGGTCCGT GACTATGGCTCCCCAGAGCCTGCCTTCATCTAGGATGGCTCCTCTGGGCATGCTGCTTGG GCTGCTGATGGCCGCCTGCTTCACCTTCTGCCTCAGTCATCAGAACCTGAAGGAGTTTGC CCTGACCAACCCAGAGAAGAGCAGCACCAAAGAAACGGAGAGAAAAGAAACCAAAGCCGA GGAGGAGCTGGATGCCGAAGTCCTGGAGGTGTTCCACCCGACGCATGAGTGGCAGGCCCT TCAGCCAGGGCAGGCTGTCCCTGCAGGATCCCACGTACGGCTGAATCTTCAGACTGGGGA AAGAGAGGCAAAACTCCAATATGAGGACAAGTTCCGAAATAATTTGAAAGGCAAAAGGCT GGATATCAACACCAACACCTACACATCTCAGGATCTCAAGAGTGCACTGGCAAAATTCAA GCTCTTCCGCCCCATTGAGGAACTGAAGAAAGACTTTGATGAGCTGAATGTTGTCATTGA GACTGACATGCAGATCATGGTACGGCTGATCAACAAGTTCAATAGTTCCAGCTCCAGTTT GGAAGAGAAGATTGCTGCGCTCTTTGATCTTGAATATTATGTCCATCAGATGGACAATGC GCAGGACCTGCTTTCCTTTGGTGGTCTTCAAGTGGTGATCAATGGGCTGAACAGCACAGA GCCCCTCGTGAAGGAGTATGCTGCGTTTGTGCTGGGCGCTGCCTTTTCCAGCAACCCCAA GGTCCAGGTGGAGGCCATCGAAGGGGGAGCCCTGCAGAAGCTGCTGGTCATCCTGGCCAC GGAGCAGCCGCTCACTGCAAAGAAGAAGGTCCTGTTTGCACTGTGCTCCCTGCTGCGCCA CTTCCCCTATGCCCAGCGGCAGTTCCTGAAGCTCGGGGGGCTGCAGGTCCTGAGGACCCT GGTGCAGGAGAAGGGCACGGAGGTGCTCGCCGTGCGCGTGGTCACACTGCTCTACGACCT GGTCACGGAGAAGATGTTCGCCGAGGAGGAGGCTGAGCTGACCCAGGAGATGTCCCCAGA GAAGCTGCAGCAGTATCGCCAGGTACACCTCCTGCCAGGCCTGTGGGAACAGGGCTGGTG CGAGATCACGGCCCACCTCCTGGCGCTGCCCGAGCATGATGCCCGTGAGAAGGTGCTGCA GACACTGGGCGTCCTCCTGACCACCTGCCGGGACCGCTACCGTCAGGACCCCCAGCTCGG CAGGACACTGGCCAGCCTGCAGGCTGAGTACCAGGTGCTGGCCAGCCTGGAGCTGCAGGA TGGTGAGGACGAGGCTACTTCCAGGAGCTGCTGGGCTCTGTCAACAGCTTGCTGAAGGA GCTGAGATGAGGCCCCACACCAGGACTGGACTGGGATGCCGCTAGTGAGGCTGAGGGGTG CCAGCGTGGGTGGGCTTCTCAGGCAGGAGGACATCTTGGCAGTGCTGGCTTGGCCATTAA

FIGURE 162

MAPQSLPSSRMAPLGMLLGLLMAACFTFCLSHQNLKEFALTNPEKSSTKETERKETKAEE ELDABVLEVPHPTHEWQALQPGQAVPAGSHVRLNLQTGERBAKLQVEDKFRNNLKGKRLD INTNTYTSQDLKSALAKFKEGAEMESSKEDKARQAEVKRLFRPIEELKKDFDELNVVIET DMQIMVRLINKFNSSSSSLEEKIAALFDLEYYVHQMDNAQDLLSFGGLQVVINGINSTEP LVKEYAAFVLGAAFSSNPKVQVEAIEGGALQKLLVILATEQPLTAKKKVLFALCSLLRHF PYAQRQFLKLGGLQVURTLVQEKGTEVLAVRVVTLLYDLVTEKMFAEEEABLTQEMSPEK LQQYRQVHLLPGLWEGWCEITAHLLALPEHDAREKVLQTIGVLLTTCRDRYRQDPQLGR TIASLOAEVQVLASLELQDGEDBGYFQELLGSVNSLIKELR

Important features: Signal peptide: amino acids 1-29

Hypothetical YJL126w/YLR351c/yhcX family protein: amino acids 364-373

N-glycosylation site: amino acids 193-197, 236-240

N-myristoylation site: amino acids 15-21, 19-25, 234-240, 251-257, 402-408, 451-457

Homologous region SLS1 protein: amino acids 68-340

FIGURE 163

CAGAGAGGAGGCTTTGGGAATTGTCCAGCAGAAACAGAGAGTCTGAGGTGGTGTCAAGA CAAAAGATGCTTCAGCTTTGGAAACTTGTTCTCCTGTGCGGCGTGCTCACTGGGACCTCA GAGTCTCTTCTTGACAATCTTGGCAATGACCTAAGCAATGTCGTGGATAAGCTGGAACCT GTTCTTCACGAGGGACTTGAGACAGTTGACAATACTCTTAAAGGCATCCTTGAGAAACTG AAGGTCGACCTAGGAGTGCTTCAGAAATCCAGTGCTTGGCAACTGGCCAAGCAGAAGGCC CAGGAAGCTGAGAAATTGCTGAACAATGTCATTTCTAAGCTGCTTCCAACTAACACGGAC GATGGCAAAGGCCTTAACCTGAGCTTCCCTGTCACCGCGAATGTCACTGTGGCCGGGCCC ATCATTGGCCAGATTATCAACCTGAAAGCCTCCTTGGACCTCCTGACCGCAGTCACAATT GAAACTGATCCCCAGACACCAGCCTGTTGCCGTCCTGGGAGAATGCGCCAGTGACCCA ACCAGCATCTCACTTTCCTTGCTGGACAAACACAGCCAAATCATCAACAAGTTCGTGAAT AGCGTGATCAACACGCTGAAAAGCACTGTATCCTCCCTGCTGCAGAAGGAGATATGTCCA CTGATCCGCATCTTCATCCACTCCCTGGATGTGAATGTCATTCAGCAGGTCGTCGATAAT CCTCAGCACAAAACCCAGCTGCAAACCCTCATCTGAAGAGGACGAATGAGGAGGACCACT GGAAGCTGCTGCCACCACCTAACCAGCGTGAAAGCCTGAGTCCCACCAGAAGGACCTTCC CAGATACCCCTTCTCCTCACAGTCAGAACAGCAGCCTCTACACATGTTGTCCTGCCCCTG GCAATAAAGGCCCATTTCTGCACCCTTAA

FIGURE 164

MLQLWKLVLLCGVLTGTSESLLDNLGNDLSNVVDKLEPVLHEGLETVDNTLKGILEKLKV DLGVLQKSSAWQLAKQKAQEAEKLLNNVISKLLPTNTDIFGLKISNSLILDVKAEFIDDG KGINLSFPVTANVTVAGFIIGQIINLKASLDLLTAVTIETDPQTHQPVAVLGECASDPTS ISLSLLDKHSQIINKFVNSVINTLKSTVSSLLQKEICPLIRIFIHSLDVNVIQQVVDNPQ HKTOLOTLI

Important features:

Signal peptide: 1-15

Transmembrane domain:

N-glycosylation site: 124-128, 132-136

N-myristoylation site: 12-18, 16-22, 26-32, 101-107, 122-128, 141-147

Leucine zipper pattern: 44-66

FIGURE 165

GCAGTCAGAGACTTCCCCTGCCCCTCGCTGGGAAAGAACATTAGGAATGCCTTTTAGTGC CTTGCTTCCTGAACTAGCTCACAGTAGCCCGGCGGCCCAGGGCAATCCGACCACATTTCA CTCTCACCGCTGTAGGAATCCAG**ATG**CAGGCCAAGTACAGCAGCACGAGGGACATGCTGG ATGATGATGGGGACACCACCATGAGCCTGCATTCTCAAGCCTCTGCCACAACTCGGCATC CAGAGCCCCGGCGCACAGAGCACAGGGCTCCCTCTTCAACGTGGCGACCAGTGGCCCTGA CCCTGCTGACTTTGTGCTTGGTGCTGCTGATAGGGCTGGCAGCCCTGGGGCTTTTGTTTT TAGGAAATACGTCCCAAGAGTTGCAATCTCTTCAAGTCCAGAATATAAAGCTTGCAGGAA GTCTGCAGCATGTGGCTGAAAAACTCTGTCGTGAGCTGTATAACAAAGCTGGAGCACACA GGTGCAGCCCTTGTACAGAACAATGGAAATGGCATGGAGACAATTGCTACCAGTTCTATA AAGACAGCAAAAGTTGGGAGGACTGTAAATATTTCTGCCTTAGTGAAAACTCTACCATGC TGAAGATAAACAAACAAGAAGACCTGGAATTTGCCGCGTCTCAGAGCTACTCTGAGTTTT TCTACTCTTATTGGACAGGGCTTTTGCGCCCTGACAGTGGCAAGGCCTGGCTGTGGATGG ATGGAACCCCTTTCACTTCTGAACTGTTCCATATTATAATAGATGTCACCAGCCCAAGAA GCAGAGACTGTGTGGCCATCCTCAATGGGATGATCTTCTCAAAGGACTGCAAAGAATTGA AGCGTTGTGTCTGTGAGAGAGGGCAGGAATGGTGAAGCCAGAGAGCCTCCATGTCCCCC $\tt CTGAAACATTAGGCGAAGGTGACTGATTCGCCCTCTGCAACTACAAATAGCAGAGTGAGC$ CAGGCGGTGCCAAAGCAAGGGCTAGTTGAGACATTGGGAAATGGAACATAATCAGGAAAG ACTATCTCTCTGACTAGTACAAAATGGGTTCTCGTGTTTCCTGTTCAGGATCACCAGCAT TTCTGAGCTTGGGTTTATGCACGTATTTAACAGTCACAAGAAGTCTTATTTACATGCCAC CAACCAACCTCAGAAACCCATAATGTCATCTGCCTTCTTGGCTTAGAGATAACTTTTAGC TCTCTTTCTCCAATGTCTAATATCACCTCCCTGTTTTCATGTCTTCCTTACACTTGGT GGAATAAGAAACTTTTTGAAGTAGAGGAAATACATTGAGGTAACATCCTTTTCTCTGACA GTCAAGTAGTCCATCAGAAATTGGCAGTCACTTCCCAGATTGTACCAGCAAATACACAAG GAATTCTTTTTGTTTGTTTCAGTTCATACTAGTCCCTTCCCAATCCATCAGTAAAGACCC CATCTGCCTTGTCCATGCCGTTTCCCAACAGGGATGTCACTTGATATGAGAATCTCAAAT CTCAATGCCTTATAAGCATTCCTTCCTGTGTCCATTAAGACTCTGATAATTGTCTCCCCT CCATAGGAATTTCTCCCAGGAAAGAAATATATCCCCATCTCCGTTTCATATCAGAACTAC CGTCCCCGATATTCCCTTCAGAGAGATTAAAGACCAGAAAAAAGTGAGCCTCTTCATCTG CACCTGTAATAGTTTCAGTTCCTATTTTCTTCCATTGACCCATATTTATACCTTTCAGGT ACTGAAGATTTAATAATAATAAATGTAAATACTGTGAAAAA

FIGURE 166

MQAKYSSTRDMLDDDGDTTMSLHSQASATTRHPEPRRTEHRAPSSTWRPVALTLLTLCLV LLIGLAALGLLFFQYYQLSNTGQDTISQMEERLGNTSQELQSLQVQNIKLAGSLQHVAEK LCRELYNKAGAHRCSPCTEQWKWHGDNCYQFYKDSKSWEDCKYFCLSENSTMLKINKQED LEFAASQSYSEFFYSYWTGLLRPDSGKAWLWMDGTPFTSELFHIIIDVTSPRSRDCVAIL NGMIFSKDCKFLKRCYCFRRAGMYKPESLHVPPETLGEGD

FIGURE 167

 $\tt CTCAGG{\color{red}{ATG}} AGGGGGGAATCTGGCCCTGGTGGGCGTTCTAATCAGCCTGGCCTTCCTGTCA$ CTGCTGCCATCTGGACATCCTCAGCCGGCTGGCGATGACGCCTGCTCTGTGCAGATCCTC GTCCTGGCCTCAAAGGGGATGCGGGAGAGAAGGGGAGACAAAGGCGCCCCCGGACGGCCT GGAAGAGTCGGCCCCACGGGAGAAAAAGGAGACATGGGGGACAAAGGACAGAAAGGCAGT GTGGGTCGTCATGGAAAAATTGGTCCCATTGGCTCTAAAGGTGAGAAAGGAGATTCCGGT CGCAAGGCCATCGGGGAGATGGACAACCAGGTCTCTCAGCTGACCAGCGAGCTCAAGTTC ATCAAGAATGCTGTCGCCGGTGTGCGCGAGACGGAGAGCAAGATCTACCTGCTGAAG GAGGAGAAGCGCTACGCGGACGCCCAGCTGTCCTGCCAGGGCCGCGGGGGCACGCTGAGC ATGCCCAAGGACGAGGCTGCCAATGGCCTGATGGCCGCATACCTGGCGCAAGCCGGCCTG GCCCGTGTCTTCATCGGCATCAACGACCTGGAGAAGGAGGGCGCCTTCGTGTACTCTGAC CACTCCCCCATGCGGACCTTCAACAAGTGGCGCAGCGGTGAGCCCAACAATGCCTACGAC GAGGAGGACTGCGTGGAGATGGTGGCCTCGGGCGGCTGGAACGACGTGGCCTGCCACACC ACCATGTACTTCATGTGTGAGTTTGACAAGGAGAACATGTGAGCCTCAGGCTGGGGCTGC CCATTGGGGGCCCCACATGTCCCTGCAGGGTTGGCAGGGACAGAGCCCAGACCATGGTGC CAGCCAGGGAGCTGTCCCTCTGTGAAGGGTGGAGGCTCACTGAGTAGAGGGCTGTTGTCT AAACTGAGAAAATGGCCTATGCTTAAGAGGAAAATGAAAGTGTTCCTGGGGTGCTGTCTC TGAAGAAGCAGAGTTTCATTACCTGTATTGTAGCCCCAATGTCATTATGTAATTATTACC CAGAATTGCTCTTCCATAAAGCTTGTGCCTTTGTCCAAGCTATACAATAAAATCTTTAAG

FIGURE 168

MRGNLALVGVLISLAFLSLLPSGHPQPAGDDACSVQILVPGLKGDAGEKGDKGAPGRPGR VGPTGEKGDMGDKGQKGSVGRHGKIGPIGSKGEKGDSGDIGPPGPNGEPGLPCECSQLRK AIGEMDNQVSQLTSELKFIKNAVAGVRETESKIYLLVKEEKRYADAQLSCQGRGGTLSMP KDEAANGLMAAYLAQAGLARVFIGINDLEKEGAFVYSDHSPMRTFNKWRSGEPNNAYDEE DCVPMVASGGWNDVACHTTMYFMCEFDKENM

FIGURE 169

AGTGACTGCAGCCTTCCTAGATCCCCTCCACTCGGTTTCTCTCTTTTGCAGGAGCACCGGC AGCACCAGTGTGTGAGGGGAGCAGGCAGCGGTCCTAGCCAGTTCCTTGATCCTGCCAGAC CACCCAGCCCCGGCACAGAGCTGCTCCACAGGCACCATGAGGATCATGCTGCTATTCAC GGAGGTGGTTCCTGGCGGGGGCCGCAGCAAGAGGGATCCAGATCTCTACCAGCTGCTCCA GAGACTCTTCAAAAGCCACTCATCTCTGGAGGGATTGCTCAAAGCCCTGAGCCAGGCTAG ACTTATGGGCAAGAGGAGCGTCCAGCCAGAGGGAAAGACAGGACCTTTCTTACCTTCAGT GAGGGTTCCTCGGCCCCTTCATCCCAATCAGCTTGGATCCACAGGAAAGTCTTCCCTGGG AACAGAGGAGCAGAGACCTTTATAAGACTCTCCTACGGATGTGAATCAAGAGAACGTCCC CAGCTTTGGCATCCTCAAGTATCCCCCGAGAGCAGAATAGGTACTCCACTTCCGGACTCC TGGACTGCATTAGGAAGACCTCTTTCCCTGTCCCAATCCCCAGGTGCGCACGCTCCTGTT ACCCTTTCTCTTCCCTGTTCTTGTAACATTCTTGTGCTTTGACTCCTTCTCCATCTTTTC TACCTGACCCTGGTGTGGAAACTGCATAGTGAATATCCCCAACCCCAATGGGCATTGACT GTAGAATACCCTAGAGTTCCTGTAGTGTCCTACATTAAAAATATAATGTCTCTCTATT AAAAAAAAAAA

FIGURE 170

MRIMLLFTAILAFSLAQSFGAVCKEPQEEVVPGGGRSKRDPDLYQLLQRLFKSHSSLEGL LKALSQASTDPKESTSPEKRDMHDFFVGLMGKRSVQPEGKTGPFLPSVRVPRPLHPNQLG STGKSSLGTEEQRPL

Important features: Signal peptide: amino acids 1-18

Tyrosine kinase phosphorylation site: amino acids 36-45

N-myristoylation site: amino acids 33-39, 59-65

Amidation site: amino acids 90-94

Leucine zipper pattern: amino acids 43-65

Tachykinin family signature: amino acids 86-92

FIGURE 171

TGGCCTCCCCAGCTTGCCAGGCACAAGGCTGAGCGGGAGGAAGCGAGAGGCATCTAAGCA GGCAGTGTTTTGCCTTCACCCCAAGTGACCATGAGAGGTGCCACGCGAGTCTCAATCATG CTCCTCCTAGTAACTGTGTCTGACTGTGCTGTGATCACAGGGGCCTGTGAGCGGGATGTC CAGTGTGGGGCAGGCACCTGCTGTGCCATCAGCCTGTGGCTTCGAGGGCTGCGGATGTGC ACCCCGCTGGGGCGGAAGGCGAGGAGTGCCACCCCGGCAGCCACAAGGTCCCCTTCTTC AGGAAACGCAAGCACCACACCTGTCCTTGCTTGCCCAACCTGCTGTGCTCCAGGTTCCCG GACGGCAGGTACCGCTCCATGGACTTGAAGAACATCAATTTTTAGGCGCGCTTGCCTGG TCTCAGGATACCCACCATCCTTTTCCTGAGCACAGCCTGGATTTTTATTTCTGCCATGAA ACCCAGCTCCCATGACTCTCCCAGTCCCTACACTGACTACCCTGATCTCTCTTGTCTAGT ACGCACATATGCACACAGGCAGACATACCTCCCATCATGACATGGTCCCCAGGCTGGCCT GAGGATGTCACAGCTTGAGGCTGTGGTGTGAAAGGTGGCCAGCCTGGTTCTCTTCCCTGC TCAGGCTGCCAGAGAGGTGGTAAATGGCAGAAAGGACATTCCCCCTCCCCTCCCCAGGTG ACCTGCTCTCTTTCCTGGGCCCTGCCCCTCTCCCCACATGTATCCCTCGGTCTGAATTAG ACATTCCTGGGCACAGGCTCTTGGGTGCATTGCTCAGAGTCCCAGGTCCTGGCCTGACCC TCAGGCCCTTCACGTGAGGTCTGTGAGGACCAATTTGTGGGTAGTTCATCTTCCCTCGAT TGGTTAACTCCTTAGTTTCAGACCACAGACTCAAGATTGGCTCTTCCCAGAGGGCAGCAG ACAGTCACCCCAAGGCAGGTGTAGGGAGCCCAGGGAGGCCAATCAGCCCCCTGAAGACTC TGGTCCCAGTCAGCCTGTGGCTTGTGGCCTGTGACCTGTGACCTTCTGCCAGAATTGTCA TGCCTCTGAGGCCCCCTCTTACCACACTTTACCAGTTAACCACTGAAGCCCCCCAATTCCC ACAGCTTTTCCATTAAAATGCAAATGGTGGTGGTTCAATCTAATCTGATATTGACATATT AGAAGGCAATTAGGGTGTTTCCTTAAACAACTCCTTTCCAAGGATCAGCCCTGAGAGCAG GTTGGTGACTTTGAGGAGGGCAGTCCTCTGTCCAGATTGGGGTGGGAGCAAGGGACAGGG AGCAGGGCAGGGGCTGAAAGGGGCACTGATTCAGACCAGGGAGGCAACTACACACCAACA TGCTGGCTTTAGAATAAAAGCACCAACTGAAAAAA

FIGURE 172

MRGATRVSIMLLLVTVSDCAVITGACERDVQCGAGTCCAISLWLRGLRMCTPLGREGEEC HPGSHKVPFFRRRKHHTCPCLPNLLCSRFPDGRYRCSMDLKNINF

Signal peptide:

Tyrosine kinase phosphorylation site: amino acids 88-95

N-myristoylation sites: amino acids 33-39, 35-41, 46-52

FIGURE 173

AGCGCCCGGGCGTCGGGGCGGTAAAAGGCCGGCAGAAGGGAGGCACTTGAGAA**ATG**TCTT TCCTCCAGGACCCAAGTTTCTTCACCATGGGGATGTGGTCCATTGGTGCAGGAGCCCTGG GGGCTGCTTGCCATTGCTGCTTGCCAACACAGACGTGTTTCTGTCCAAGCCCCAGA AAGCGGCCCTGGAGTACCTGGAGGATATAGACCTGAAAACACTGGAGAAGGAACCAAGGA CTTTCAAAGCAAAGGAGCTATGGGAAAAAAATGGAGCTGTGATTATGGCCGTGCGGAGGC CAGGCTGTTTCCTCTGTCGAGAGGAAGCTGCGGATCTGTCCTCCCTGAAAAGCATGTTGG ACCAGCTGGGCGTCCCCCTCTATGCAGTGGTAAAGGAGCACATCAGGACTGAAGTGAAGG ATTTCCAGCCTTATTTCAAAGGAGAAATCTTCCTGGATGAAAAGAAAAAGTTCTATGGTC CACAAAGGCGGAAGATGATGTTTATGGGATTTATCCGTCTGGGAGTGTGGTACAACTTCT TCCGAGCCTGGAACGGAGGCTTCTCTGGAAACCTGGAAGGAGAAGGCTTCATCCTTGGGG GAGTTTTCGTGGTGGGATCAGGAAAGCAGGGCATTCTTCTTGAGCACCGAGAAAAAGAAT TTGGAGACAAAGTAAACCTACTTTCTGTTCTGGAAGCTGCTAAGATGATCAAACCACAGA CTTTGGCCTCAGAGAAAAATGATTGTGTGAAACTGCCCAGCTCAGGGATAACCAGGGAC ATTCACCTGTGTTCATGGGATGTATTGTTTCCACTCGTGTCCCTAAGGAGTGAGAAACCC ATTTATACTCTACTCTCAGTATGGATTATTAATGTATTTTAATATTCTGTTTAGGCCCAC TAAGGCAAAATAGCCCCAAAACAAGACTGACAAAAATCTGAAAAACTAATGAGGATTATT AAGCTAAAACCTGGGAAATAGGAGGCTTAAAATTGACTGCCAGGCTGGGTGCAGTGGCTC ACACCTGTAATCCCAGCACTTTGGGAGGCCAAGGTGAGCAAGTCACTTGAGGTCGGGAGT TCGAGACCAGCCTGAGCAACATGGCGAAACCCCGTCTCTACTAAAAATACAAAAATCACC CGGGTGTGGCAGCACCTGTAGTCCCAGCTACCCGGGAGGCTGAGGCAGGAGAATCA CTTGAACCTGGGAGGTGGAGGTTGCGGTGAGCTGAGATCACACCACTGTATTCCAGCCTG GGTGACTGAGACTCTAACTAA

FIGURE 174

MSFLQDPSFFTMGMWSIGAGALGAAALALLLANTDVFLSKPQKAALEYLEDIDLKTLEKE PRTFKAKELWEKNGAVIMAVRRPGCFLCREEAADLSSLKSMLDQLGVPLYAVVKEHIRTE VKDFQPYFKGEIFLDEKKKFYGPQRRKMMFMGFIRLGVWYNFFRAWNGGFSGNLEGEGFI LGGVPVVGSGKQGILLEHREKEFEGDKVNLLSVLEAAKMIKPQTLASEKK

FIGURE 175

GACAGTGGAGGGCAGTGGAGAGGACCGCGCTGTCCTGCTGTCACCAAGAGCTGGAGACAC CATCTCCCACCGAGAGTC<u>ATG</u>GCCCCATTGGCCCTGCACCTCCTCGTCCTCGTCCCCATC CTCCTCAGCCTGGTGGCCTCCCAGGACTGGAAGGCTGAACGCAGCCAAGACCCCTTCGAG AAATGCATGCAGGATCCTGACTATGAGCAGCTGCTCAAGGTGGTGACCTGGGGGCTCAAT CGGACCCTGAAGCCCCAGAGGGTGATTGTGGTTGGCGCTGGTGTGGCCGGGCTGGTGGCC GCCAAGGTGCTCAGCGATGCTGGACACAAGGTCACCATCCTGGAGGCAGATAACAGGATC GGGGGCCGCATCTTCACCTACCGGGACCAGAACACGGGCTGGATTGGGGAGCTGGGAGCC ATGCGCATGCCCAGCTCTCACAGGATCCTCCACAAGCTCTGCCAGGGCCTGGGGCTCAAC CTGACCAAGTTCACCCAGTACGACAAGAACACGTGGACGGAGGTGCACGAAGTGAAGCTG CGCAACTATGTGGTGGAGAAGGTGCCCGAGAAGCTGGGCTACGCCTTGCGTCCCCAGGAA AAGGGCCACTCGCCCGAAGACATCTACCAGATGGCTCTCAACCAGGCCCTCAAAGACCTC AAGGCACTGGGCTGCAGAAAGGCGATGAAGAAGTTTGAAAGGCACACGCTCTTGGAATAT CTTCTCGGGGAGGGGAACCTGAGCCGGCCGGCCGTGCAGCTTCTGGGAGACGTGATGTCC GAGGATGGCTTCTTCTATCTCAGCTTCGCCGAGGCCCTCCGGGCCCACAGCTGCCTCAGC AGCTCGCTGTCCGGGCTTGTGCTGTTGAACGCGCCCGTGGTGGCGATGACCCAGGGACCG CACGATGTGCACGTGCAGATCGAGACCTCTCCCCCGGCGCGGAATCTGAAGGTGCTGAAG GCCGACGTGGTGCTGCTGACGGCGAGCGGACCGGCGGTGAAGCGCATCACCTTCTCGCCG CCGCTGCCCCGCCACATGCAGGAGGCGCTGCGGAGGCTGCACTACGTGCCGGCCACCAAG GTGTTCCTAAGCTTCCGCAGGCCCTTCTGGCGCGAGGAGCACATTGAAGGCGGCCACTCA AACACCGATCGCCCGTCGCGCATGATTTTCTACCCGCCGCCGCGCGAGGGCGCGCTGCTG CTGGCCTCGTACACGTGGTCGGACGCGGCGGCAGCGTTCGCCGGCTTGAGCCGGGAAGAG GCGTTGCGCTTGGCGCTCGACGACGTGGCGGCATTGCACGGGCCTGTCGTGCGCCAGCTC TGGGACGGCACCGGCGTCGTCAAGCGTTGGGCGGAGGACCAGCACAGCCAGGGTGGCTTT GTGGTACAGCCGCCGCGCTCTGGCAAACCGAAAAGGATGACTGGACGGTCCCTTATGGC CGCATCTACTTTGCCGGCGAGCACACCGCCTACCCGCACGGCTGGGTGGAGACGGCGGTC AAGTCGGCGCTGCGCCCCCATCAAGATCAACAGCCGGAAGGGGCCTGCATCGGACACG GCCAGCCCGAGGGGCACGCATCTGACATGGAGGGGCAGGGGCATGTGCATGGGGTGGCC AGCAGCCCCTCGCATGACCTGGCAAAGGAAGAAGGCAGCCACCCTCCAGTCCAAGGCCAG TTATCTCTCCAAAACACGACCCACACGAGGACCTCGCAT<u>TAA</u>AGTATTTTCGGAAAAAAA

FIGURE 176

MAPLALHLLVLVPILLSLVASQDWKAERSQDPFEKCMQDPDYEQLLKVVTWGLNRTLKPQ
RVIVVGAGVAGLVAAKVLSDAGHKVTILEADNRIGGRIFTTRDQNTGWIGBLGAMRMPSHRILHKLCQGLGLNLTKFTQYDKNTWTEVHEVKLRNYVVEKVPEKLGYALRPQEKGHSPE
DIYQMALNQALKDLKALGCRKAMKKFERHTLLEYLLGEGNLSRPAVQLLGDVMSEDGFFY
LSFAEALRAHSCLSDRLQYSRIVGGWDLLPRALLSSLSGLVLLNAPVVAMTQGPHDVHVV
IETSPPARNLKVLKADVVLLTASGPAVKRITFSPPLPRHMQEALRRLHYVPATKVFLSFR
RPFWREEHIEGGHSNTDRPSRMIFYPPPREGALLLASYTWSDAAAAFAGLSREEALRLAL
DDVAALHGPVVRQLWDGTGVVKRWAEDQHSQGGFVVQPPALWQTEKDDWTVPYGRIYFAG
EHTAYPHGWVETAVKSALRAAIKINSRKGPASDTASPEGHASDMEGQGHVHGVASSPSHD
LAKEEGSHPPVQGQLSLQNTTHTRTSH

Signal peptide: amino acids 1-21

FIGURE 177

CCGGGGAGGGGAGGGCCCGTCCCGCCCCTCCCCGTCTCTCCCCGCCCCTCCCCGTCCCTC CCGCCGAAGCTCCGTCCCGCCGCGGGCCGGCTCCGCCCTCACCTCCCGGCCGCGGCTGC CTGCTGTTCGTGGCGCTACCCGCCTCCGGCTGGCTGACGACGGCCCCCCGAGCCGCCG CCGCTGTCCGGAGCCCCACAGGACGGCATCAGAATTAATGTAACTACACTGAAAGATGAT GGGGACATATCTAAACAGCAGGTTGTTCTTAACATAACCTATGAGAGTGGACAGGTGTAT GTAAATGACTTACCTGTAAATAGTGGTGTAACCCGAATAAGCTGTCAGACTTTGATAGTG AAGAATGAAAATCTTGAAAATTTGGAGGAAAAAGAATATTTTGGAATTGTCAGTGTAAGG ATTTTAGTTCATGAGTGGCCTATGACATCTGGTTCCAGTTTGCAACTAATTGTCATTCAA GAAGAGGTAGTAGAGATTGATGGAAAACAAGTTCAGCAAAAGGATGTCACTGAAATTGAT AAAAAAGAAAGTGTTAGTTCACTGCAAACCACTAGCCAGTATCTTATCAGGAATGTGGAA ACCACTGTAGATGAAGATGTTTTACCTGGCAAGTTACCTGAAACTCCTCTCAGAGCAGAG CCGCCATCTTCATATAAGGTAATGTGTCAGTGGATGGAAAAGTTTAGAAAAGATCTGTGT AGGTTCTGGAGCAACGTTTTCCCAGTATTCTTTCAGTTTTTGAACATCATGGTGGTTGGA ATTACAGGAGCAGCTGTGGTAATAACCATCTTAAAGGTGTTTTTCCCAGTTTCTGAATAC AAAGGAATTCTTCAGTTGGATAAAGTGGACGTCATACCTGTGACAGCTATCAACTTATAT CCAGATGGTCCAGAGAAAAGAGCTGAAAACCTTGAAGATAAAACATGTATTTAAAACGCC ATCTCATATCATGGACTCCGAAGTAGCCTGTTGCCTCCAAATTTGCCACTTGAATATAAT TTTCTTTAAATCGTT

FIGURE 178

MEGAPPGSLALRLLLFVALPASGWLTTGAPEPPPLSGAPQDGIRINVTTLKDDGDISKQQ
VVLNITYESGQVYVNDLPVNSGVTRISCQTLIVKNENLENLEEKEYEGIVSVRILVHEW
MTSGSSLQLIVIQEEVVEIDGKQVQKDVTEIDILVKNRGVLRHSNTTPLEESMLYSIS
RDSDILFTLPNLSKKESVSSLQTTSQYLIRNVETTVDEDVLPGKLPETPLRAEPPSSYKV
MCQMMEKFRKDLCRFWSNVFPVFFQFLNIMVVGITGAAVVITILKVFFPVSEYKGILQLD
KVDVIPVTAINLYPDGPEKRAENLEDKTCI

Signal peptide:

Transmembrane domain: 266-284

Leucine zipper pattern: 155-177

N-glycosylation site: 46-50, 64-68, 166-170, 191-195

Motif name: N-myristoylation site: 3-9, 42-48, 273-279

FIGURE 179

CTCCTTAGGTGGAAACCCTGGGAGTAGAGTACTGACAGCAAAGACCGGGAAAGACCATAC GTCCCCGGGCAGGGGTGACAACAGGTGTCATCTTTTTGATCTCGTGTGTGGGCTGCCTTCC TATTTCAAGGAAAGACGCCAAGGTAATTTTGACCCAGAGGAGCAATGATGTAGCCACCTC CTAACCTTCCCTTCTTGAACCCCCAGTTATGCCAGGATTTACTAGAGAGTGTCAACTCAA CCAGCAAGCGGCTCCTTCGGCTTAACTTGTGGTTGGAGGAGAACCTTTGTGGGGCTGC GTTCTCTTAGCAGTGCTCAGAAGTGACTTGCCTGAGGGTGGACCAGAAGAAAGGAAAGGT CCCCTCTTGCTCGCTCCACACACAGGAAGGCTGTGATGGGAATGAAGGTGAAACTTG GAGATTTCACTTCAGTCATTGCTTCTGCCTGCAAGATCATCCTTTAAAAGTAGAGAAGCT GCTCTGTGTGGTGGTTAACTCCAAGAGGCAGAACTCGTTCTAGAAGGAAATGGATGCAAG CAGCTCCGGGGGCCCCAAACGCATGCTTCCTGTGGTCTAGCCCAGGGAAGCCCTTCCGTG GGGGCCCGGCTTTGAGGGATGCCACCGGTTCTGGACGCATGGCTGATTCCTGAATGATG ATGGTTCGCCGGGGGCTGCTTGCGTGGATTTCCCGGGTGGTGGTTTTTGCTGGTGCTCCTC TGCTGTGCTATCTCTCTCTCTGTACATGTTGGCCTGCACCCCAAAAGGTGACGAGGAGCAG CTGGCACTGCCCAGGGCCAACAGCCCCACGGGGAAGGAGGGGTACCAGGCCGTCCTTCAG GAGTGGGAGGAGCAGCACCGCAACTACGTGAGCAGCCTGAAGCGGCAGATCGCACAGCTC AAGGAGGAGCTGCAGGAGGAGGAGCTGAGCAGCTCAGGAATGGGCAGTACCAAGCCAGCGAT GCTGCTGGCCTGGGTCTGGACAGGAGCCCCCCAGAGAAAACCCAGGCCGACCTCCTGGCC TTCCTGCACTCGCAGGTGGACAAGGCAGAGGTGAATGCTGGCGTCAAGCTGGCCACAGAG TATGCAGCAGTGCCTTTCGATAGCTTTACTCTACAGAAGGTGTACCAGCTGGAGACTGGC CTTACCCGCCACCCCGAGGAGAGCCTGTGAGGAAGGACAAGCCGGGATGAGTTGGTGGAA GCCATTGAATCAGCCTTGGAGACCCTGAACAATCCTGCAGAGAACAGCCCCAATCACCGT CCTTACACGGCCTCTGATTTCATAGAAGGGATCTACCGAACAGAAAGGGACAAAGGGACA TTGTATGAGCTCACCTTCAAAGGGGACCACAAACACGAATTCAAACGGCTCATCTTATTT CGACCATTCAGCCCCATCATGAAAGTGAAAAATGAAAAGCTCAACATGGCCAACACGCTT ATCAATGTTATCGTGCCTCTAGCAAAAAGGGTGGACAAGTTCCGGCAGTTCATGCAGAAT TTCAGGGAGATGTGCATTGAGCAGGATGGGAGAGTCCATCTCACTGTTGTTTACTTTGGG AAAGAAGAAATAAATGAAGTCAAAGGAATACTTGAAAACACTTCCAAAGCTGCCAACTTC AGGAACTTTACCTTCATCCAGCTGAATGGAGAATTTTCTCGGGGAAAGGGACTTGATGTT GGAGCCCGCTTCTGGAAGGGAAGCAACGTCCTTCTCTTTTTCTGTGATGTGGACATCTAC TTCACATCTGAATTCCTCAATACGTGTAGGCTGAATACACAGCCAGGGAAGAAGGTATTT TATCCAGTTCTTTTCAGTCAGTACAATCCTGGCATAATATACGGCCACCATGATGCAGTC CCTCCCTTGGAACAGCAGCTGGTCATAAAGAAGGAAACTGGATTTTGGAGAGACTTTGGA TTTGGGATGACGTGTCAGTATCGGTCAGACTTCATCAATATAGGTGGGTTTGATCTGGAC ATCA A AGGCTGGGGCGGAGAGGATGTGCACCTTTATCGCAAGTATCTCCACAGCAACCTC ATAGTGGTACGGACGCCTGTGCGAGGACTCTTCCACCTCTGGCATGAGAAGCGCTGCATG GACGAGCTGACCCCGAGCAGTACAAGATGTGCATGCAGTCCAAGGCCATGAACGAGGCA TCCCACGGCCAGCTGGGCATGCTGGTGTTCAGGCACGAGATAGAGGCTCACCTTCGCAAA CAGAAACAGAAGACAAGTAGCAAAAAAACATGAACTCCCAGAGAAGGATTGTGGGAGACA CTTTTTCTTTCCTTTTGCAATTACTGAAAGTGGCTGCAACAGAGAAAAGACTTCCATAAA GGACGACAAAAGAATTGGACTGATGGGTCAGAGATGAGAAAGCCTCCGATTTCTCTCTGT TGGGCTTTTTACAACAGAAATCAAAATCTCCGCTTTGCCTGCAAAAGTAACCCAGTTGCA CCCTGTGAAGTGTCTGACAAAGGCAGAATGCTTGTGAGATTATAAGCCTAATGGTGTGGA GGTTTTGATGGTGTTTACAATACACTGAGACCTGTTGTTTTTGTGTGCTCATTGAAATATT CATGATTTAAGAGCAGTTTTGTAAAAAATTCATTAGCATGAAAGGCAAGCATATTTCTCC TCATATGAATGAGCCTATCAGCAGGGCTCTAGTTTCTAGGAATGCTAAAATATCAGAAGG CAGGAGAGAGATAGGCTTATTATGATACTAGTGAGTACATTAAGTAAAATAAAATGGAC CAGAAAAGAAAGAAACCATAAATATCGTGTCATATTTTCCCCAAGATTAACCAAAAATA

 $\mathsf{TGCACTTTTTTCCCTTGTGAGTTATAGTCTGCTTATTTAATTACCACTTTGCAAGCCTT$ AATACAGACGTACAGATACTTTCTCTGAAGAGTATTTTCGAAGAGGAGCAACTGAACACT GGAGGAAAAGAAAATGACACTTTCTGCTTTACAGAAAAGGAAACTCATTCAGACTGGTGA TATCGTGATGTACCTAAAAGTCAGAAACCACATTTTCTCCTCAGAAGTAGGGACCGCTTT CTTACCTGTTTAAATAAACCAAAGTATACCGTGTGAACCAAACAATCTCTTTTCAAAACA TATATATATTCTCAAACATCAATCCATCTCCCAGAATCTAGTGGGATGGAAGTTTTTTGCT ACATGTTATCCACCCCAGGCCAGGTGGAAGTAACTGAATTATTTTTAAATTAAGCAGTT CTACTCAATCACCAAGATGCTTCTGAAAATTCCATTTTATTACCATTTCAAACTATTTTTTCACCAGATGCATGAGCTAATTATCTCTTTGAGTCCTTGCTTCTGTTTGCTCACAGTAAAC TCATTGTTTAAAAGCTTCAAGAACATTCAAGCTGTTGGTGTGTTAAAAAAATGCATTGTAT TGATTTGTACTGGTAGTTTATGAAATTTAATTAAAACACAGGCCATGAATGGAAGGTGGT ATTGCACAGCTAATAAAATATGATTTGTGGATATGAA

FIGURE 180

MMMVRRGLLAWISRVVVLLVLLCCAISVLYMLACTPKGDEEQLALPRANSPTGKEGYQAV
LQEWEEQHRNYVSSLKRQIAQLKEELQERSEQLRNGQYQASDAGLGLDRSPPEKTQADL
LAFLHSQVDKAEVNAGVKLATE¥XAVPPDSFTLQKVYQLETGLTRHFEEKPVKKDKRDEL
VEAIESALETLNNPAENSPNHRPYTASDFIEGIYRTERDKGTLYELTFKGDHKHEFKRLI
LFRPFSPIMKVKNEKLNMANTLINVIVPLAKRVDKFRQFMQNFREMCIEQDGRVHLTVVY
FGKEEINEVKGILENTSKAANFRNFTFIQLNGEFSRGKGLDVGARFWKGSNVLLFFCDVD
IYFTSEFLNTCRLNTQPGKKVFYPVLFSQYNFGIIYGHHDAVPPLEQQLVIKKETGFWRD
FGFGMTCQYRSDFINIGGFDLDIKGWGGEDVHLYRKYLHSNLIVVRTPVRGLFHLWHEKR
CMDELTPEGYKMCMQSKAMNEASHGQLGMLVFRHEIEAHLRKQKQKTSSKKT

FIGURE 181

 $\tt CGTCTCTGCGTTCGCC{\color{red}{\textbf{ATG}}} CGTCCCGGGGCGCCAGGGCCACTCTGGCCTCTGCCCTGGGG$ GGCCTGGCTTGGGCCGTGGGCTTCGTGAGCTCCATGGGCTCGGGGAACCCCGCGCCCCGG TGGTGTTTGCTGGCTCCAGCAGGGCCAGGAGGCCACCTGCAGCCTGGTGCTCCAGACTGA TGTCACCCGGGCCGAGTGCTGTGCCTCCGGCAACATTGACACCGCCTGGTCCAACCTCAC CCACCCGGGGAACAAGATCAACCTCCTCGGCTTCTTGGGCCTTGTCCACTGCCTTCCCTG CAAAGATTCGTGCGACGCGTGGAGTGCGGCCCGGGCAAGGCGTGCCGCATGCTGGGGGG CCACCCGGACCTGAGCGTCATGTACCGGGGCCGCTGCCGCAAGTCCTGTGAGCACGTGGT GTGCCCGCGGCCACAGTCGTGCGTCGTGGACCAGACGGGCAGCGCCCACTGCGTGGTGTG TCGAGCGGCCCTGCCCTGTGCCCTCCAGCCCCGGCCAGGAGCTTTGCGGCAACAACAA CGTCACCTACATCTCCTCGTGCCACATGCGCCAGGCCACCTGCTTCCTGGGCCGCTCCAT CGGCGTGCGCCACGCGGGCAGCTGCGCAGGCACCCCTGAGGAGCCGCCAGGTGGTGAGTC TGCAGAAGAGAAGAGAACTTCGTGTGAGCCTGCAGGACAGGCCTGGGCCTGGTGCCCGA GGCCCCCATCATCCCCTGTTATTTATTGCCACAGCAGAGTCTAATTTATATGCCACGGA CACTCCTTAGAGCCCGGATTCGGACCACTTGGGGATCCCAGAACCTCCCTGACGATATCC CGGACACTGAGCGCCTGATTTAGGGCCCTTCTCTAGGATGCCCCAGCCCCTACCCTAAGA CTACTATCAAGAGGGCTGGGCATTCTCTGCTGGTAATTCCTGAAGAGGCATGACTGCTTT TCTCAGCCCCAAGCCTCTAGTCTGGGTGTGTACGGAGGGTCTAGCCTGGGTGTGTACGGA GGGTCTAGCCTGGGTGAGTACGGAGGGTCTAGCCTGGGTGAGTACGGAGGGTCTAGCCTG GGTGAGTACGGAGGGTCTAGCCTGGGTGTGTATGGAGGATCTAGCCTGGGTGAGTATGGA GGGTCTAGCCTGGGTGAGTATGGAGGGTCTAGCCTGGGTGTGTATGGAGGGTCTAGCCTG GGTGAGTATGGAGGGTCTAGCCTGGGTGTGTATGGAGGGTCTAGCCTGGGTGAGTATGGA GGGTCTAGCCTGGGTGTGTACGGAGGGTCTAGTCTGAGTGCGTGTGGGGACCTCAGAACA CTGTGACCTTAGCCCAGCAAGCCAGGCCCTTCATGAAGGCCAAGAAGGCTGCCACCATTC CCTGCCAGCCCAAGAACTCCAGCTTCCCCACTGCCTCTGTGTGCCCCTTTGCGTCCTGTG AAGGCCATTGAGAAATGCCCAGTGTGCCCCCTGGGAAAGGGCACGGCCTGTGCTCCTGAC ACGGGCTGTGCTTGGCCACAGAACCACCCAGCGTCTCCCCTGCTGCTGTCCACGTCAGTT ACTGTGTCCGGCGGAGCCAAGTCCACTCTGGGGGAGCTCTGGCGGGGACCACGGGCCACT GCTCACCCACTGGCCCCGAGGGGGGTGTAGACGCCAAGACTCACGCATGTGTGACATCCG GAGTCCTGGAGCCGGGTGTCCCAGTGGCACCACTAGGTGCCTGCTGCCTCCACAGTGGGG TTCACACCCAGGGCTCCTTGGTCCCCCACAACCTGCCCCGGCCAGGCCTGCAGACCCAGA TGGTCTTGGGCCAGTTCTCCCACGACGGCTCACCCTCCCCTCCATCTGCGTTGATGCTCA GAATCGCCTACCTGTGCCTGCGTGTAAACCACAGCCTCAGACCAGCTATGGGGAGAGGAC AACACGGAGGATATCCAGCTTCCCCGGTCTGGGGTGAGGAATGTGGGGAGCTTGGGCATC CCTAGGTTGGTGGGTCTACAGGAGCCTCAGCCAGGCAGCCCACCCCACCCTGGGGCCCTG CCTCACCAAGGAAATAAAGACTCAAGCCATAAAAAAAA

FIGURE 182

MRPGAPGPLWPLPWGALAWAVGFVSSMGSGNPAPGGVCWLQQGQEATCSLVLQTDVTRAE CCASGNIDTAWSNLTHPGNKINLLGFLGLVHCLPCKDSCDGVECGFGKACKMLGGRPRCE CAPDCSGLPARLQVCGSDGATYRDECELRAARCRGHPDLSVMYRGRCRKSCEHVVCPRPQ SCVVDQTGSAHCVVCRAAPCPVPSSPGQELCGNNNVTYISSCHMRQATCFLGRSIGVRHA GSCAGTPEEPPGGESAEEEENFV

Important features: Signal peptide: amino acids 1-20

N-glycosylation sites: amino acids 73-77, 215-219

Osteonectin domain proteins: amino acids 97-130, 169-202

FIGURE 183

CACTCATTCATTCCAAAGGGTCTCTCAAGGCAATGGTAATGTGCAAGGAGGTGATACCTA AATGAATGACCAAAAGAACATGCTTCTGCTTTTGTGTGTCTCCTACATTTTAGACATTTG TTTGTTTCTCTTGGTAGCCTTTAAATTCCTTGAAGCCCAGGACCATGTCTCACTTACCTT TGTGTTTCCACTAACTAGTCTACCTCCTGGAATTGGCAGATACTCAGTGAAAGCCTGTGA AATAAGTGATGTCTATTTCTAGCATATTATTCTGAGATTTAATGATAGATTTAGTGATTG TTCAACTTCATTCTCAAAATTAGGTCCTGAGTTAACTAATAATTACCTTTGAAATGTGTG TGATTCAGTTCCATTGCATTGATTTTTGTTCTCAGAAGCCAAGGTTTCCCATGAAAAATC ATTCCCACTTGAATTGGGCTGTGATTCTTGCTGCGTTTAAGTAAAGGAAGCCTCTTGGTT CTAGTTCTGCAAACTTACACACTGAACTGGGACAAGTTTTTGTTTAGAGTAATGGCTGGG AAAAGAGGAACCTTTCATTTTATTCAGAAGTCAAAAACAAAGGCCTCCCAGCCACCTGGA GATGTTTTGTTGCAGACACCAGCCTGGCTCTGTCTTTATGCCTAACAATTGAGCATCCAG TCTTCTTTGTGCTGGGACCATTGCTCAGCTCTGCAAGGGGAAAAGAGGGAGAAAGCCAGA GCTGCCAGGCTTCTTGCACTGGGGCCGGGGGAGGGTTCCTGGGAAGCAGGTGCTCTCTGG TTTGCTCCAGGAGTATATTGAAAACATACCCAGTGCTCTCTCAAGCACCCACTGCTTAGA GGGCCCAGATTTCTTTTCCTTCTTTCCCTTGCAGAGCTGGAGACTGCATCGGGCATCTGG TGTTTAAACTAAACAGGAAAACTGACTAAAGGTCCACAGTGCTCATTGTGTAGACTAGCT GCCCTCCGATGGGTGCTCTGATTATCAGTGGTTCCAGTGCAGGGCCTGTCACTAAACAGG CCTCACTTCCTCCTTGGGGGCTTTCCCATGGGAGGTGTGGCTTTTTACTCTACATGGAAA TGACTCTCTGCAGCCACAGAACACAGTCATTTTCTGAATTATCCCAGTCTCTCATGCGCC CTGGATTCCTCCAGATGCCTTATATCTCTTGTGCAAAGTTGTCTAAAATTTGGTTCCCAG CTTCCAAGCCTTGCCTTTTGGCCTTCCTGGAAGTATTTTTGTTGATGAGTCGTCTGTCAT TATTCTCTAAAATGATTTGCTTTTTGTTTCTTTCATTCCTATTTCCACCCCACATATACA CACATGCTTCTTAACTTAGGGGATTACATGCCAATAAATCTATTGTTGAAAATGCACTAA TACTATCGCAAAGACGAAAATTCACAGGCTGAACCGTTGTAAGTCCATATGCTCCTCAAC TTACATGTGTGATGGAGTTATGCCCAAATAAGTCCATCGTCAAGTTGAAAAATCAAAATC AAGCCATCTTAGGTTGAGGACCATTTGTTTGTACCTCCAAAGATGTCATATCTTTAAACA TACTCCCTAGCTTTTCTTTTACTTTTATTTTGAAGTAATTATAGAATCACAGAAAGTT GCAAAAAA

FIGURE 184

MGALIISGSSAGPVTKQASLPPWGLSHGRCGFLLYMEMTLCSHRTQSFSELSQSLMRPGF LQMPYISCAKLSKIWFPASKPCLLAFLEVFLLMSRLSLFSKMICFLFLSFLFPPHIYTHAS

Important features of the protein: Signal peptide: amino acids 1-41

Transmembrane domain: amino acids 88-107

Casein kinase II phosphorylation site: amino acids 47-50

N-myristoylation site: amino acids 24-29

FIGURE 185

AACTCAAACTCCTCTCTGGGAAAACGCGGTGCTTGCTCCTCCCGGAGTGGCCTTGGCA GGGTGTTGGAGCCCTCGGTCTGCCCCGTCCGGTCTCTGGGGCCAAGGCTGGGTTTCCCTC ATGTATGGCAAGAGCTCTACTCGTGCGGTGCTTCTTCTCCTTGGCATACAGCTCACAGCT CTTTGGCCTATAGCAGCTGTGGAAATTTATACCTCCCGGGTGCTGGAGGCTGTTAATGGG ACAGATGCTCGGTTAAAATGCACTTTCTCCAGCTTTGCCCCTGTGGGTGATGCTCTAACA $\tt GTGACCTGGAATTTTCGTCCTCTAGACGGGGGACCTGAGCAGTTTGTATTCTACTACCAC$ ATAGATCCCTTCCAACCCATGAGTGGGCGCTTTTAAGGACCGGGTGTCTTGGGATGGGAAT CCTGAGCGGTACGATGCCTCCATCCTTCTCTGGAAACTGCAGTTCGACGACAATGGGACA TACACCTGCCAGGTGAAGAACCCACCTGATGTTGATGGGGTGATAGGGGAGATCCGGCTC AGCGTCGTGCACACTGTACGCTTCTCTGAGATCCACTTCCTGGCTCTGGCCATTGGCTCT GCCTGTGCACTGATGATCATAATAGTAATTGTAGTGGTCCTCTTCCAGCATTACCGGAAA AAGCGATGGGCCGAAAGAGCTCATAAAGTGGTGGAGATAAAATCAAAAGAAGAGGAAAGG CTCAACCAAGAGAAAAAGGTCTCTGTTTATTTAGAAGACACAGACTAACAATTTTAGATG TTCTTTGGCTTTTCCAGTTGTGACCCGTTTTCCAACCAGTTCTGCAGCATATTAGATTCT AGACAAGCAACACCCCTCTGGAGCCAGCACAGTGCTCCTCCATATCACCAGTCATACACA GCCTCATTATTA AGGTCTTATTTAATTTCAGAGTGTAAATTTTTTCAAGTGCTCATTAGG TTTTATAAACAAGAAGCTACATTTTTGCCCTTAAGACACTACTTACAGTGTTATGACTTG TATACACATATATTCGTATCAAACGCGATAAAACCCAATTTCTCTCTTACATTTCCTTTC ACGTATTTCTTTTAGCAGCACTTCTGCTACTAAAGTTAATGTGTTTACTCTTTCCTTC CCACATTCTCAATTAAAAGGTGAGCTAAGCCTCCTCGGTGTTTCTGATTAACAGTAAATC CTAAATTCAAACTGTTAAATGACATTTTTATTTTTATGTCTCTCCTTAACTATGAGACAC ATCTTGTTTTACTGAATTTCTTTCAATATTCCAGGTGATAGATTTTTGTCG

FIGURE 186

MYGKSSTRAVLLLLGIQLTALWPIAAVEIYTSRVLEAVNGTDARLKCTFSSFAPVGDALT VTWMFRPLDGGPEGFVFYYHIDPFQPMSGRFKDRVSWDGNPERYDASILLWKLQFDDNGT YTCQVKNPPDVDGVIGEIRLSVVHTVRFSEIHFLALAIGSACALMIIIVIVVVLFQHYRK KRWAERAHKVVEIKSKEEERLNQEKKVSVYLEDTD

FIGURE 187

FIGURE 188

 ${\tt MKFLAVLVLLGVSIFLVSAQNPTTAAPADTYPATGPADDEAPDAETTAAATTATTAAPTT} \\ {\tt ATTAASTTARKDIPVLPKWVGDLPNGRVCP}$

FIGURE 189

GAGCGAACATGGCAGCGCGTTGGCGGTTTTGGTGTGTCTCTGTGACCATGGTGGTGGCGC GAGACAAGTTCCGTCGCCTTGTGAAAGCCCCACCGAGAAATTACTCCGTTATCGTCATGT AGATCCTGGCAAACTCCTGGCGATACTCCAGTGCATTCACCAACAGGATATTTTTTGCCA TGGTGGATTTTGATGAAGGCTCTGATGTATTTCAGATGCTAAACATGAATTCAGCTCCAA CTTTCATCAACTTTCCTGCAAAAGGGAAACCCAAACGGGGTGATACATATGAGTTACAGG TGCGGGGTTTTTCAGCTGAGCAGATTGCCCGGTGGATCGCCGACAGAACTGATGTCAATA TTAGAGTGATTAGACCCCCAAATTATGCTGGTCCCCTTATGTTGGGATTGCTTTTGGCTG TTATTGGTGGACTTGTGTATCTTCGAAGAAGTAATATGGAATTTCTCTTTAATAAAACTG GATGGGCTTTTGCAGCTTTGTGTTTTGTGCTTGCTATGACATCTGGTCAAATGTGGAACC ATATAAGAGGACCACCATATGCCCATAAGAATCCCCACACGGGACATGTGAATTATATCC ATGGAAGCAGTCAAGCCCAGTTTGTAGCTGAAACACACATTGTTCTTCTGTTTAATGGTG GAGTTACCTTAGGAATGGTGCTTTTATGTGAAGCTGCTACCTCTGACATGGATATTGGAA AGCGAAAGATAATGTGTGGGCTGGTATTGGACTTGTTGTATTATTCTTCAGTTGGATGC TCTCTATTTTTAGATCTAAATATCATGGCTACCCATACAGCTTTCTGATGAGT**TAA**AAAG TGAAAAGAAGAATGCAACTTGTATATTTTGTATTACCTCTTTTTTTCAAGTGATTTAAAT AGTTAATCATTTAACCAAAGAAGATGTGTAGTGCCTTAACAAGCAATCCTCTGTCAAAAT ATTTAATTTAGTACAATTAAGTATATTATAAAAATTGTAAAACTACTACTTTGTTTTAGT TAGAACAAAGCTCAAAACTACTTTAGTTAACTTGGTCATCTGATTTTATATTGCCTTATC CAAAGATGGGGAAAGTAAGTCCTGACCAGGTGTTCCCACATATGCCTGTTACAGATAACT ACATTAGGAATTCATTCTTAGCTTCTTCATCTTTGTGTGGATGTGTATACTTTACGCATC TTTCCTTTTGAGTAGAGAAATTATGTGTGTCATGTGGTCTTCTGAAAATGGAACACCATT CTTCAGAGCACACGTCTAGCCCTCAGCAAGACAGTTGTTTCTCCTCCTCCTTGCATATTT AATACAGGATTATAATTTCTGCTTGAGTATGGTGTTAACTACCTTGTATTTAGAAAGATT TCAGATTCATTCCATCTCCTTAGTTTTCTTTTAAGGTGACCCATCTGTGATAAAAATATA GCTTAGTGCTAAAATCAGTGTAACTTATACATGGCCTAAAATGTTTCTACAAATTAGAGT TTGTCACTTATTCCATTTGTACCTAAGAGAAAAATAGGCTCAGTTAGAAAAGGACTCCCT TCACGAGGTCAGGAGTTCGAGACCATCCTGGCCAACATGGTGAAACCCCGTCTCTACTAA AAATATAAAAATTAGCTGGGTGTGGTGGCAGGAGCCTGTAATCCCAGCTACACAGGAGGC TGAGGCACGAGAATCACTTGAACTCAGGAGATGGAGGTTTCAGTGAGCCGAGATCACGCC

FIGURE 190

MAARWFFWCYSVTMYVALLIVCDVPSASAQRKKEMVLSEKVSQLMEWTNKRPVIRMNGDK FRRLVKAPPRNYSVIVMFTALQLHRQCVVCKQADEEFQILANSWRYSSAFTNRIFFAMVK FDEGSDVFCMLNNNSAPTFINFPAKGKPKRGDTYELQVRGFSAEQIARWIADRTDVNIRV IRPPNYAGPLMLGLLLAVIGGLVYLRRSNMEFLFNKTGWAFAALCFVLAMTSQQMWNHIR GPPYAHKNPHTGHVNYIHGSSQAQFVAETHIVLLFNGGVTLGMVLLCEAATSDMDIGKRK IMCVAGIGLUVULFFSWMLSIFRSKYHGYPYSFLMS

Signal peptide: amino acids 1-29

Transmembrane domains: amino acids 183-205, 217-237, 217-287, 301-321

FIGURE 191

GAGAGAAGTCAGCCTGGCAGAGAGACTCTGAAATGAGGGATTAGAGGTGTTCAAGGAGCA AGAGCTTCAGCCTGAAGACAAGGGAGCAGTCCCTGAAGACGCTTCTACTGAGAGGTCTGC CATGGCCTCTCTTGGCCTCCAACTTGTGGGCTACATCCTAGGCCTTCTGGGGCTTTTGGG CACACTGGTTGCCATGCTGCTCCCCAGCTGGAAAACAAGTTCTTATGTCGGTGCCAGCAT CATCACCCAGTGTGACATCTATAGCACCCTTCTGGGCCTGCCCGCTGACATCCAGGCTGC CCAGGCCATGATGGTGACATCCAGTGCAATCTCCTCCCTGGCCTGCATTATCTCTGTGGT GGGCATGAGATGCACAGTCTTCTGCCAGGAATCCCGAGCCAAAGACAGAGTGGCGGTAGC TCATGGGATCCTACGGGACTTCTACTCACCACTGGTGCCTGACAGCATGAAATTTGAGAT TGGAGAGGCTCTTTACTTGGGCATTATTTCTTCCCTGTTCTCCCTGATAGCTGGAATCAT CCTCTGCTTTTCCTGCTCATCCCAGAGAAATCGCTCCAACTACTACGATGCCTACCAAGC CCAACCTCTTGCCACAAGGAGCTCTCCAAGGCCTGGTCAACCTCCCAAAGTCAAGAGTGA GTTCAATTCCTACAGCCTGACAGGGTATGTG**TGA**AGAACCAGGGGCCAGAGCTGGGGGGT GGCTGGGTCTGTGAAAAACAGTGGACAGCACCCCGAGGGCCACAGGTGAGGGACACTACC AAGGCAGAAATGGGGGCTAGTGTAACAGCATGCAGGTTGAATTGCCAAGGATGCTCGCCA TGCCAGCCTTTCTGTTTTCCTCACCTTGCTGCTCCCCTGCCCTAAGTCCCCAACCCTCAA CTTGAAACCCCATTCCCTTAAGCCAGGACTCAGAGGATCCCTTTGCCCTCTGGTTTACCT GGCTTTTGTGGGCATTGCTCTAACCTACTTCTCAAGCTTCCCTCCAAAGAAACTGATTGG CCCTGGAACCTCCATCCCACTCTTGTTATGACTCCACAGTGTCCAGACTAATTTGTGCAT GAACTGAAATAAAACCATCCTACGGTATCCAGGGAACAGAAAGCAGGATGCAGGATGGGA GGACAGGAAGGCAGCCTGGGACATTTAAAAAAATA

FIGURE 192

MASLGLQLVGYILGLLGLLGTLVAMLLPSWKTSSYVGASIVTAVGFSKGLWMECATHSTG ITQCDIYSTLLGLPADIQAAQAMMVTSSAISSLACIISVVGMRCTVFCQESRAKDRVAVA GGVFFILGGLLGFIPVAWNLHGILRDFYSPLVPDSMKFEIGEALYLGIISSLFSLIAGII LCFSCSSQRNRSNYYDAYQAQPLATRSSPRFGQPPKVKSEFNSYSLTGYV

Important features of the protein: Signal peptide: amino acids 1-24

Transmembrane domains: amino acids 82-102, 117-140, 163-182

N-glycosylation site: amino acids 190-193

PMP-22 / EMP / MP20 family proteins: amino acids 46-59

FIGURE 193

CTCCACTGCAACCACAGAGCCATGGCTCCCCGAGGCTGCATCGTAGCTGTTTTGCCA
TTTTCTGCATCTCCAGGCTCCTCTGCTCACACGGAGCCCCATGATCCCCATGACTCCTT
ACCTGATGCTGTGCCAGCCACAAGAGATGTGGGGACAAGTGTACGACCCCTGCAG
ACTGTTGCTATGATGATGCCGTCGTGCCCTTGGCCAGGACCTATGACACCTCCAGACCTTTCAGACTCCTTTCAGACTCCTTGCCAGGACCTCAGACCTGTGAAAACTGCA
CCTTCAGAGTCTGCTTTGAGCAGTGCCCCTGGACCTTCATGGTGAAAGCTGATAAAC
AGAACTGCGACTCAGCCCGGACCTCGGATGACAGGCCTTCATGGTGCAGTGTCAGCTAATGGA
ACATCAGGGGAACCATGACTCCTGGATTCTCCTTCCTGGGTGGCCTGGAGAAAGACGCT
GGTGTTACCTGAGATCTGGATGCTGAGTGGCTTTTGGGGCCAGAGAAACACACTC
AACTGCCCACTTCATTCTGTGACCTGTCTGAGGCCCACCTGCCCTGAGGAGGC
CCACAGGTCCCCTTCTAGAATTCTGGACAGCATCAGATCGTTCTCTTCTGAGAGGCCCAG
GGACTCTGAACCCTCCTGATGACCCCTATGGCCAACATCAACCCGGCACCACCCCAAGGC
TGGCTGGGGAACCTTCACCCTTCTGTGAGATTTTCCATCATCTCAAGTTCTCTTCTATC
CAGGAGCAAAGCACAGGATCATAATAAAATTTATGTACTTTATAAATGAAAA

FIGURE 194

MAPRGCIVAVFAIFCISRLLCSHGAPVAPMTPYLMLCQPHKRCGDKFYDPLQHCCYDDAV VPLARTQTCGNCTFRVCFEQCCPWTFMVKLINQNCDSARTSDDRLCRSVS

Signal peptide: amino acids 1-24

FIGURE 195

FIGURE 196

MMLHSALGLCLLLVTVSSNLAIAIKKEKRPPQTLSRGWGDDITWVQTYEEGLFYAQKSKK PLMVIHHLEDCQYSQALKKVFAQNEEIQEMAQNKFIMLNLMHETTDKNLSPDGQYVPRIM FVDPSLTVRADIAGRYSNRLYTYEPRDLPLLIENMKKALRLIQSEL

Important features: Signal peptide: amino acids 1-23

N-myristoylation site: amino acids 51-57

FIGURE 197

AGTCGCGGAGGCTTCCCCGCGCCGGCCGCGCCCGCTCCCCGGCACCAGAAGTTC CTCTGCGCGTCCGACGCGACATGGGCGTCCCCACGGCCCTGGAGGCCGGCAGCTGGCGC TGGGGATCCCTGCTCTTCGCTCTCTTCCTGGCTGCGTCCCTAGGTCCGGTGGCAGCCTTC AAGGTCGCCACGCCGTATTCCCTGTATGTCTGTCCCGAGGGGCAGAACGTCACCCTCACC TGCAGGCTCTTGGGCCCTGTGGACAAAGGGCACGATGTGACCTTCTACAAGACGTGGTAC CGCAGCTCGAGGGGCGAGGTGCAGACCTGCTCAGAGCGCCGGCCCATCCGCAACCTCACG TTCCAGGACCTTCACCTGCACCATGGAGGCCACCAGGCTGCCAACACCAGCCACGACCTG GCTCAGCGCCACGGGCTGGAGTCGGCCTCCGACCACCATGGCAACTTCTCCATCACCATG CGCAACCTGACCCTGCTGGATAGCGGCCTCTACTGCTGCCTGGTGGTGGAGATCAGGCAC CACCACTCGGAGCACAGGGTCCATGGTGCCATGGAGCTGCAGGTGCAGACAGGCAAAGAT GCACCATCCAACTGTGTGTGTACCCATCCTCCTCCCAGGATAGTGAAAACATCACGGCT GCAGCCCTGGCTACGGGTGCCTGCATCGTAGGAATCCTCTGCCTCCCCCTCATCCTGCTC CTGGTCTACAAGCAAAGGCAGGCAGCCTCCAACCGCCGTGCCCAGGAGCTGGTGCGGATG GACAGCAACATTCAAGGGATTGAAAACCCCGGCTTTGAAGCCTCACCACCTGCCCAGGGG ATACCCGAGGCCAAAGTCAGGCACCCCCTGTCCTATGTGGCCCAGCGGCAGCCTTCTGAG TCTGGGCGGCATCTGCTTTCGGAGCCCAGCACCCCCCTGTCTCCCAGGCCCCGGAGAC GTCTTCTTCCCATCCTGGACCCTGTCCCTGACTCTCCAAACTTTGAGGTCATCTAGCCC AGCTGGGGGACAGTGGGCTGTTGTGGCTGGGTCTGGGGCAGGTGCATTTGAGCCAGGGCT GATACTGTGACATCCCAGAAGCCCAGCCCCTCAACCCCTCTGGATGCTACATGGGGATGC TGGACGGCTCAGCCCCTGTTCCAAGGATTTTGGGGTGCTGAGATTCTCCCCTAGAGACCT GAAATTCACCAGCTACAGATGCCAAATGACTTACATCTTAAGAAGTCTCAGAACGTCCAG CCCTTCAGCAGCTCTCGTTCTGAGACATGAGCCTTGGGATGTGGCAGCATCAGTGGGACA AGATGGACACTGGGCCACCTCCCAGGCACCAGACACAGGGCACGGTGGAGAGACTTCTC CCCCGTGGCCGCCTTGGCTCCCCGTTTTGCCCGAGGCTGCTCTTCTGTCAGACTTCCTC TTTCTACCACAGTGGCTCTGGGGCCAGGCCTGCCTGCCCACTGGCCATCGCCACCTTCCC CAGCTGCCTCCTACCAGCAGTTTCTCTGAAGATCTGTCAACAGGTTAAGTCAATCTGGGG CTTCCACTGCCTGCATTCCAGTCCCCAGAGCTTGGTGGTCCCGAAACGGGAAGTACATAT TGGGGCATGGTGGCCTCCGTGAGCAAATGGTGTCTTGGGCAATCTGAGGCCAGGACAGAT GTGGAGAGGGGCACCTGCCCCGCCCTCCCCATCCCCTACTCCCACTGCTCAGCGCGGG CCATTGCAAGGGTGCCACAATGTCTTGTCCACCCTGGGACACTTCTGAGTATGAAGCG AAGA

FIGURE 198

MGVPTALEAGSWRWGSLLFALFLAASLGPVAAFKVATPYSLYVCPEGQNVTLTCRLLGPV DKGHDVTFYKTWYRSSRGEVQTCSERRPIRNLTTQDLHLHHGGHQAANTSHDLAQRHGLE SASDHHGHFSITMRNLTLLDSGLYCCLVVEIRHHHSEHRVHGAMELQVQTGKDAPSNCVV YPSSSQDSENITAAALATGACIVGILCLPLILLLVYKQRQAASNRRAQELVRMDSNIQGI ENPGFEASPPAQGIPEAKVRHPLSYVAQRQPSESGRHLLSEPSTPLSPPGPGDVFFPSLD PVPDSPNPEVI

Signal peptide: amino acids 1-28

Transmembrane domain: amino acids 190-216

FIGURE 199

FIGURE 200

MDSLRKMLISVAMLGAGAGVGYALLVIVTPGERRKQEMLKEMPLQDPRSREEAARTQQLLLATLQEAATTQENVAWRKNWMVGGEGGASGRSP

Signal peptide: amino acids 1-18

FIGURE 201

GACAGCTGTGTCTCGATGGAGTAGACTCTCAGAACAGCGCAGTTTGCCCTCCGCTCACGC AGAGCCTCTCCGTGGCTTCCGCACCTTGAGCATTAGGCCAGTTCTCCTCTTCTCTAAT CCATCCGTCACCTCTCTGTCATCCGTTTCCATGCCGTGAGGTCCATTCACAGAACACAT CCATGGCTCTCATGCTCAGTTTTGGTTCTGAGTCTCCTCAAGCTGGGATCAGGGCAGTGGCAGCTCTTTGGGCCAGACAAGCCTGTCCAGGCCTTGGTGGGGGAGGACGCAGCATTCTCCT GTTTCCTGTCTCCTAAGACCAATGCAGAGGCCATGGAAGTGCGGTTCTTCAGGGGCCAGT TCTCTAGCGTGGTCCACCTCTACAGGGACGGGAAGGACCAGCCATTTATGCAGATGCCAC AGTATCAAGGCAGGACAAAACTGGTGAAGGATTCTATTGCGGAGGGGCGCATCTCTCTGA GGCTGGAAAACATTACTGTGTTGGATGCTGGCCTCTATGGGTGCAGGATTAGTTCCCAGT TTTCCATCACGGGATATGTTGATAGAGACATCCAGCTACTCTGTCAGTCCTCGGGCTGGT GGACAAACAGAGACATGCATGGCCTGTTTGATGTGGAGATCTCTCTGACCGTCCAAGAGA ACGCCGGGAGCATATCCTGTTCCATGCGGCATGCTCATCTGAGCCGAGAGGTGGAATCCA GGGTACAGATAGGAGATACCTTTTTCGAGCCTATATCGTGGCACCTGGCTACCAAAGTAC ${\tt TGGGAATACTCTGCTGTGGCCTATTTTTTGGCATTGTTGGACTGAAGATTTTCTTCTCCA}$ AATTCCAGTGGAAAATCCAGGCGGAACTGGACTGGAGAAGAAAGCACGGACAGGCAGAAT TGAGAGACGCCCGGAAACACGCAGTGGAGGTGACTCTGGATCCAGAGACGCTCACCCGA AGCTCTGCGTTTCTGATCTGAAAACTGTAACCCATAGAAAAGCTCCCCAGGAGGTGCCTC ACTCTGAGAAGAGATTTACAAGGAAGAGTGTGGTGGCTTCTCAGAGTTTCCAAGCAGGGA AACATTACTGGGAGGTGGACGGAGGACACAATAAAAGGTGGCGCGTGGGAGTGTGCCGGG ATGATGTGGACAGGAGGAAGGAGTACGTGACTTTGTCTCCCGATCATGGGTACTGGGTCC TCAGACTGAATGGAGAACATTTGTATTTCACATTAAATCCCCGTTTTATCAGCGTCTTCC CCAGGACCCCACCTACAAAAATAGGGGTCTTCCTGGACTATGAGTGTGGGACCATCTCCT TCTTCAACATAAATGACCAGTCCCTTATTTATACCCTGACATGTCGGTTTGAAGGCTTAT TGAGGCCCTACATTGAGTATCCGTCCTATAATGAGCAAAATGGAACTCCCATAGTCATCT GCCCAGTCACCCAGGAATCAGAGAAAAGAGGCCTCTTGGCAAAGGGCCTCTGCAATCCCAG AGACAAGCAACAGTGAGTCCTCCTCACAGGCAACCACGCCCTTCCTCCCCAGGGGTGAAA CAAAGTCCCGCAGCAGCCGGCCAAGGTGGCTTCCAGATGAAGGGGGACTGGCCTGTCCAC AGTTTGCTCTCACTCCATCTGGCTAAGTGATCTTGAAATACCACCTCTCAGGTGAAGAAC CGTCAGGAATTCCCATCTCACAGGCTGTGGTGTAGATTAAGTAGACAAGGAATGTGAATA ATGCTTAGATCTTATTGATGACAGAGTGTATCCTAATGGTTTGTTCATTATATTACACTT TCAGTAAAAAA

FIGURE 202

MALMLSLVLSLLKLGSGQWQVFGPDKPVQALVGEDAAFSCFLSPKTNAEAMEVRFFRGQF SSVVHLYRDGKDQFFMMMPQYQGRTKLVKDSIAEGRISLRLENITVLDAGLYGCRISSQ YQKAIWELQVSALGSVPLISITGYVDRDIQLLCQSSGWFPRPTAKKWRGQQGDLSTDSR TNRDMHGLFDVEISLTVQENAGSISCSMRHAHLSREVESRVQIGDTFFEPISWHLATKVL GILCCGLFFGIVGLKIFFSKFQWKIQAELDWRRKHGQAELRDARKHAVEVTLDPETAHPK LCVSDLKTVTHRKAPQEVPHSEKRFTKRSVVASQSFQAGKHYWEVDGGHNKRWRVGVCRD DVDRRKEYVTLSPDHGYWVLRLNGEHLYFTLNPRFISVFPRTPPTKIGVFLDYECGTISF FNINDQSLIYTLTCRFEGLLRPYIEYPSYNEQNGTPIVICPVTQESEKEASWQRASAIPE TSNSESSSOATTFPLPRGEM

Signal peptide: amino acids 1-17

Transmembrane domain: amino acids 239-255

FIGURE 203

FIGURE 204

MGGLLLAAFLALVSVPRAQAVWLGRLDPEQLLGPWYVLAVASREKGFAMEKDMKNVVGVV VTLTPENNLRTLSSQHGLGGCDQSVMDLIKRNSGWVFENPSIGVLELWVLATNFRDYAII FTOLEFGBPFNTVELYSLTETASQEAMGLFTKWSRSLGFLSQ

Signal peptide: amino acids 1-20

FIGURE 205

FIGURE 206

MKTLFLGVTLGLAAALSFTLEEEDITGTWYVKAMVVDKDFPEDRRPRKVSPVKVTALGGG KLEATFTFMREDRCIQKKILMRKTEEPGKYSAYGGRKLMYLQELPRRDHYIFYCKDQHHG GLLHMGKLVGRNSDTNREALEEFKKLVQRKGLSEEDIFTPLQTGSCVPEH

Important features: Signal peptide: amino acids 1-17

FIGURE 207

GTTCCGCAGATGCAGAGGTTGAGGTGGCTGCGGGACTGGAAGTCATCGGGCAGAGGTCTC ACAGCAGCCAAGGAACCTGGGGCCCGCTCCTCCCCCCTCCAGGCCATGAGGATTCTGCAG TTAATCCTGCTTGCTCTGGCAACAGGGCTTGTAGGGGGAGAGACCAGGATCATCAAGGGG TTCGAGTGCAAGCCTCACTCCCAGCCCTGGCAGGCAGCCCTGTTCGAGAAGACGCGGCTA CTCTGTGGGGCGACGCTCATCGCCCCCAGATGGCTCCTGACAGCAGCCCACTGCCTCAAG CCCCGCTACATAGTTCACCTGGGGCAGCACCACCTCCAGAAGGAGGAGGGCTGTGAGCAG ACCCGGACAGCCACTGAGTCCTTCCCCCACCCGGCTTCAACAACAGCCTCCCCAACAAA GACCACCGCAATGACATCATGCTGGTGAAGATGGCATCGCCAGTCTCCATCACCTGGGCT GTGCGACCCTCACCCTCTCACGCTGTGTCACTGCTGGCACCAGCTGCCTCATTTCC GGCTGGGGCAGCACGTCCAGCCCCCAGTTACGCCTGCCTCACACCTTGCGATGCGCCAAC ATCACCATCATTGAGCACCAGAAGTGTGAGAACGCCTACCCCGGCAACATCACAGACACC ATGGTGTGTGCCAGCGTGCAGGAAGGGGGCAAGGACTCCTGCCAGGGTGACTCCGGGGGC CCTCTGGTCTGTAACCAGTCTCTTCAAGGCATTATCTCCTGGGGCCAGGATCCGTGTGCG ATCACCCGAAAGCCTGGTGTCTACACGAAAGTCTGCAAATATGTGGACTGGATCCAGGAG ACGATGAAGAACAATTAGACTGGACCCACCCACCACCACCCATCACCCTCCATTTCCACTTGGTGTTTGGTTCCTGTTCACTCTGTTAATAAGAAACCCTAAGCCAAGACCCTCTACGAA CATTCTTTGGGCCTCCTGGACTACAGGAGATGCTGTCACTTAATAATCAACCTGGGGTTC GAAATCAGTGAGACCTGGATTCAAATTCTGCCTTGAAATATTGTGACTCTGGGAATGACA ACACCTGGTTTGTTCTCTGTTGTATCCCCAGCCCCAAAGACAGCTCCTGGCCATATATCA AAAA

FIGURE 208

MRILQLILLALATGLVGGETRIIKGFECKPHSQPWQAALFEKTRLLCGATLIAPRWLLTA AHCLKPRYIVHLGQHNLQKEEGCEQTRTATESFPHPGFNNSLPNKDHRNDIMLVKMASPV SITWAVRPLTLSSRCVTAGTSCLISGWGSTSSPQLRLPHTLRCANITIIEHQKCENAYPG NITDTMVCASVQEGGKDSCQGDSGGPLVCNQSLQGIISWGQDPCAITRKPGVYTKVCKYV DWIOETMKNN

Important features: Signal peptide: amino acids 1-18

Serine proteases, trypsin family, histidine active site: amino acids 58-63

N-glycosylation sites: amino acids 99-102, 165-168, 181-184, 210-213

Glycosaminoglycan attachment site: amino acids 145-148

Kringle domain proteins: amino acids 197-209, 47-64

Serine proteases, trypsin family, histidine protein: amino acids 199-209, 47-63, 220-243

Apple domain proteins: amino acids 222-249, 189-222

2.18/392

FIGURE 209

GCGGCCACACGCAGCTAGCCGGAGCCGGACCAGGCGCCTGTGCCTCCTCCTCGTCCCTC GCCGCGTCCGCGAGCCTGGAGCCGGCGGGAGCCCCGCGCTCGCCATGTCGGGCGAGCTC AGCAACAGGTTCCAAGGAGGGAAGGCGTTCGGCTTGCTCAAAGCCCGGCAGGAGAGGAGG CTGGCCGAGATCAACCGGGAGTTTCTGTGTGACCAGAAGTACAGTGATGAAGAGAACCTT CCAGAAAAGCTCACAGCCTTCAAAGAGAAGTACATGGAGTTTGACCTGAACAATGAAGGC GAGATTGACCTGATGTCTTTTAAAGAGGATGATGAGAGAAGCTTGGTGTCCCCAAGACCCAC CTGGAGATGAGAAGATGATCTCAGAGGTGACAGGAGGGGTCAGTGACACTATATCCTAC CGAGACTTTGTGAACATGATGCTGGGGAAACGGTCGGCTGTCCTCAAGTTAGTCATGATG ATTGCTAGCCTGCCCTGAGGACCCCGCCTGGACTCCCCAGCCTTCCCACCCCATACCTCC CATTGAGGGTTTGTTTGTGTTTTCATCAATGTCTTTGTAAAGCACAAATTATCTGCCTTA TCCCGCTCCCTGTGCAGAAGGGCTGATATCAAACCAAAAACTAGAGGGGGCAGGGCCAG GGCAGGGAGGCTTCCAGCTGTGTTCCCCTCACTTGGAGGAACCAGCACTCTCCATCCTT TCAGAAAGTCTCCAAGCCAAGTTCAGGCTCACTGACCTGGCTCTGACGAGGACCCCAGGC CACTCTGAGAAGACCTTGGAGTAGGGACAAGGCTGCAGGGCCTCTTTCGGGTTTCCTTGG ACAGTGCCATGGTTCCAGTGCTCTGGTGTCACCCAGGACACAGCCACTCGGGGCCCCGCT GCCCAGCTGATCCCCACTCATTCCACACCTCTTCTCATCCTCAGTGATGTGAAGGTGGG AAGGAAAGGAGCTTGGCATTGGGAGCCCTTCAAGAAGGTACCAGAAGGAACCCTCCAGTC $\tt CTGCTCTCTGGCCACACCTGTGCAGGCAGCTGAGAGGCAGCGTGCAGCCCTACTGTCCCT$ TACTGGGGCAGCAGAGGGCTTCGGAGGCAGAAGTGAGGCCTGGGGTTTGGGGGGAAAGGT CAGCTCAGTGCTGTTCCACCTTTTAGGGAGGGTACTGAGGGGACCAGGATGGGAGAATGA GGAGTAAAATGCTCACGGCAAAGTCAGCAGCACTGGTAAGCCAAGACTGAGAAATACAAG

FIGURE 210

MSGELSNRFQGGKAFGLLKARQERRLAEINREFLCDQKYSDEENLPEKLTAFKEKYMEFD LNNEGEIDLMSLKRMMEKLGVPKTHLEMKKMISEVTGGVSDTISYRDFVNMMLGKRSAVL KLVMMFEGKANESSPKPVGPPPERDIASLP

FIGURE 211

FIGURE 212

MRRLLLVTSLVVVLLWEAGAVPAPKVPIKMQVKHWPSEQDPEKAWGARVVEPPEKDDQLV VLFPVQKPKLLTTEEKPRGGGROFILBGTKAWMETEDTLGRVLSPEPDHDSLYHPPPEED OGEERPRLWWMPHHOVLLGPEEDDHTYHPO

FIGURE 213

CAGGCAGAAGCGAACAAAGACCCAGCAAGAGAAGGCAGAGGCTAAGACCCATCCCGTATC TGCTCTCCTGAAATAATTCTGGAGTCATGCCTGAAATGCCAGAGGACATGGAGCAGGAGG AAGTTAACATCCCTAATAGGAGGGTTCTGGTTACTGGTGCCACTGGGCTTCTTGGCAGAG CTGTACACAAAGAATTTCAGCAGAATAATTGGCATGCAGTTGGCTGTGGTTTCAGAAGAG CAAGACCAAAATTTGAACAGGTTAATCTGTTGGATTCTAATGCAGTTCATCACATCATTC AAAATCAGCCAGATGCTGCCTCTCAACTTAATGTGGATGCTTCTGGGAATTTAGCAAAGG AAGCAGCTGCTGTTGGAGCATTTCTCATCTACATTAGCTCAGATTATGTATTTGATGGAA CAAATCCACCTTACAGAGAGGAAGACATACCAGCTCCCCTAAATTTGTATGGCAAAACAA AATTAGATGGAGAAAAGGCTGTCCTGGAGAACAATCTAGGAGCTGCTGTTTTGAGGATTC CTATTCTGTATGGGGAAGTTGAAAAGCTCGAAGAAAGTGCTGTGACTGTTATGTTTGATA AAGTGCAGTTCAGCAACAAGTCAGCAAACATGGATCACTGGCAGCAGAGGTTCCCCACAC ATGTCAAAGATGTGGCCACTGTGTGCCGGCAGCTAGCAGAAGAAGAATGCTGGATCCAT CAATTAAGGGAACCTTTCACTGGTCTGGCAATGAACAGATGACTAAGTATGAAATGGCAT GTGCAATTGCAGATGCCTTCAACCTCCCAGCAGTCACTTAAGACCTATTACTGACAGCC CTGTCCTAGGAGCACAACGTCCGAGAAATGCTCAGCTTGACTGCTCCAAATTGGAGACCT TGGGCATTGGCCAACGAACACCATTTCGAATTGGAATCAAAGAATCACTTTGGCCTTTCC TCATTGACAAGAGATGGAGACAAACGGTCTTTCATTAGTTTATTTGTGTTGGGTTCTTTT TTTTTTTAAATGAAAAGTATAGTATGTGGCACTTTTTAAAGAACAAAGGAAATAGTTTTG TATGAGTACTTTAATTGTGACTCTTAGGATCTTTCAGGTAAATGATGCTCTTGCACTAGT GAAATTGTCTAAAGAAACTAAAGGGCAGTCATGCCCTGTTTGCAGTAATTTTTCTTTTTA TCATTTTGTTTGTCCTGGCTAAACTTGGAGTTTGAGTATAGTAAATTATGATCCTTAAAT ATTTGAGAGTCAGGATGAAGCAGATCTGCTGTAGACTTTTCAGATGAAATTGTTCATTCT CGTAACCTCCATATTTTCAGGATTTTTGAAGCTGTTGACCTTTTCATGTTGATTATTTTA AATTGTGTGAAATAGTATAAAAATCATTGGTGTTCATTATTTGCTTTGCCTGAGCTCAGA TCAAAATGTTTGAAGAAAGGAACTTTATTTTTGCAAGTTACGTACAGTTTTTATGCTTGA GATATTTCAACATGTTATGTATATTGGAACTTCTACAGCTTGATGCCTCCTGCTTTTATA TTGAATGCAAACGTGTATTTTTTAATATAAATATAACTGTCCTTTTCATCCCATGTT GCCGCTAAGTGATATTTCATATGTGTGGTTATACTCATAATAATGGGCCTTGTAAGTCTT TTCACCATTCATGAATAATAATAATATGTACTGCTGGCATGTAATGCTTAGTTTTCTTG TATTTACTTCTTTTTTTAAATGTAAGGACCAAACTTCTAAACTAATTGTTCTTTTGTTGC TTTAATATGTATTGAAATAAAACACAATAAAATT

FIGURE 214

MPEMPEDMEQEEVNIPNRVLVTGATGLIGRAVHKEFQQNNWHAVGGGFRARPKFEQVN LLDSNAVHHIIHDFQPHVIVHCAAERRPDVVENQPDAASQLNVDASGNLAKEAAAVGAFL IYISSDYVPGGTNPPYREEDIPAPLNLYGKTKLDGEKAVLENNLGAAVLRIPILYGEVEK LEESAVTVMFDKVQFSNKSANMDHWQQRFPTHVKDVATVCRQLAEKRNLDPSIKGTFHWS GNEQMTKYEMACAIADAFNLPSSHLRPITDSFVLGAQRPRNAQLDCSKLETLGIGQRTPF RIGIKESLWFFLIDKKMROTVFH

Signal peptide:

Transmembrane domain:

N-glycosylation site: amino acids 197-201

N-myristoylation site: amino acids 303-309

Short-chain dehydrogenases/reductases family proteins: amino acids 18-30

FIGURE 215

GTGAATGTGAGGGTTTGATGACTTTCAGATGTCTAGGAACCAGAGTGGGTGCAGGGGCCC $\tt CAGGCAGGCTGATTCTTGGGCGGAGGAGGAGTAGGGTAAAGGGTTCTGCATGAGCTCCTT$ AAAGGACAAAGGTAACAGAGCCAGCGAGAGAGCTCGAGGGGAGACTTTGACTTCAAGCCA CAGAATTGGTGGAAGTGTGCGCGCCGCCGCCGCCGTCGCTCCTGCAGCGCTGTCGACCTA GCCGCTAGCATCTTCCCGAGCACCGGGATCCCGGGGTAGGAGGCGACGCGGGCGAGCACC AGCGCCAGCCGGCTGCGGCTGCCCACACGGCTCACCATGGGCTCCGGGCGCCGGGCGCTG AACGACACGGAGCCCATCGTGCTGGAGGGCAAGTGTCTGGTGGTGTGCGACTCGAACCCG GCCACGGACTCCAAGGGCTCCTCTTCCTCCCCGCTGGGGATATCGGTCCGGGCGGCCAAC TCCAAGGTCGCCTTCTCGGCGGTGCGGAGCACCAACCACGAGCCATCCGAGATGAGCAAC AAGACGCGCATCATTTACTTCGATCAGATCCTGGTGAATGTGGGTAATTTTTTCACATTG GAGTCTGTCTTTGTAGCACCAAGAAAAGGAATTTACAGTTTCAGTTTTCACGTGATTAAA GTCTACCAGAGCCAAACTATCCAGGTTAACTTGATGTTAAATGGAAAACCAGTAATATCT GCCTTTGCGGGGGACAAAGATGTTACTCGTGAAGCTGCCACGAATGGTGTCCTGCTCTAC CAGTATTCCACGTTTTCTGGCTTTCTGGTGTTCCCCCTATAGGATTCAATTTCTCCATGA TGTTCATCCAGGTGAGGGATGACCCACTCCTGAGTTATTGGAAGATCATTTTTTCATCAT TGGATTGATGTCTTTTATTGGTTTCTCATGGGTGGATATGGATTCTAAGGATTCTAGCCT TTGGGACTCTAAGCAGATAATACCTATGCTTAAATGTAACAGTCAAAAGCTGTCTGCAAG ACTTATTCTGAATTTCATTTCCTGGGATTACTGAATTAGTTACAGATGTGGAATTTTATT TGTTTAGTTTTAAAAGACTGGCAACCAGGTCTAAGGATTAGAAAACTCTAAAGTTCTGAC TTCAATCAACGGTTAGTGTGATACTGCCAAAGAACTGTATACTGTGTTAATATATTGATT ATATTTGTTTTTATTCCTTTGGAATTAGTTTGTTTGGTTCTTGTAAAAAACTTGGATTTT TAGTTGTATTTTAATTGTATATGTGAAAGAGTCATATTTTCCAAGTTATATTTTCTAAGA AGAAGAATAGATCATAAATCTGACAAGGAAAAAGTTGCTTACCCAAAATCTAAGTGCTCA ATCCCTGAGCCTCAGCAAAACAGCTCCCCTCCGAGGGAAATCTTATACTTTATTGCTCAA CGTAGACATGACCACTTTATTAACTGGTGGTGGGATGCTGTTGTTTCTAATTATACCTAT TTTTCAAGGCTTCTGTTGTATTTGAAGTATCATCTGGTTTTGCCTTAACTCTTTAAATTG TATATATTTATCTGTTTAGCTAATATTAAATTCAAATATCCCATATCTAAATTTAGTGCA TAAAGATTAATATATGTTAAAAAAA

FIGURE 216

MGSGRRALSAVPAVLLVLTLPGLPVWAQNDTEPIVLEGKCLVVCDSNPATDSKGSSSSPL GISVRAANSKVAFSAVRSTNHEPSEMSNKTRIIYFDQILVNVGNFFTLESVFVAPRKGIY SFSFHVIKVYQSQTIQVNLMLNGKPVISAFAGDKDVTREAATNGVLLYLDKEDKVYLKLE KGNLVGGWQYSTFSGFLVFPL

Signal peptide: amino acids 1-27

FIGURE 217

TGGGCTTGCCCTTCTTGGTGCTCTTGGTGGCCTCGGTCGAGAGCCATCTGGGGGTTCTGG GGCCCAAGAACGTCTCGCAGAAAGACGCCGAGTTTGAGCGCACCTACGTGGACGACGTCA ACAGCGAGCTGGTCAACATCTACACCTTCAACCATACTGTGACCCGCAACAGGACAGAGG GCGTGCGTGTCTCTGAACGTCCTGAACAAGCAGAAGGGGGCGCCGTTGCTGTTTGTGG TCCGCCAGAAGGAGGCTGTGGTGTCCTTCCAGGTGCCCCTAATCCTGCGAGGGATGTTTC AGCGCAAGTACCTCTACCAAAAAGTGGAACGAACCCTGTGTCAGCCCCCCACCAAGAATG ACCAGCTCCGGGTCAGCCGCATGGACGATTTTGTGCTCAGGACTGGGGAGCAGTTCAGCT TCAATACCACAGCAGCACCCCAGTACTTCAAGTATGAGTTCCCTGAAGGCGTGGACT CGCTA ATTGTCA AGGTGA CCTCCA ACA AGGCCTTCCCCTGCTCAGTCATCTCCATTCAGG ATGTGCTGTGTCCTGTCTATGACCTGGACAACAACGTAGCCTTCATCGGCATGTACCAGA CGATGACCAAGAAGGCGGCCATCACCGTACAGCGCAAAGACTTCCCCAGCAACAGCTTTT ATGTGGTGGTGGTGAAGACCGAAGACCAAGCCTGCGGGGGCTCCCTGCCTTTCTACC CCTTCGCAGAGATGAACCGGTCGATCAAGGGCACCGCCAGAAAACCCTGTCAGTGCTGG TGTCTCAAGCAGTCACGTCTGAGGCATACGTCAGTGGGATGCTCTTTTGCCTGGGTATAT TTCTCTCTTTTACCTGCTGACCGTCCTCCTGGCCTGCTGGGAGAACTGGAGGCAGAAGA AGAAGACCCTGCTGGTGGCCATTGACCGAGCCTGCCCAGAAAGCGGTCACCCTCGAGTCC TGGCTGATTCTTTTCCTGGCAGTTCCCCTTATGAGGGTTACAACTATGGCTCCTTTGAGA ATGTTTCTGGATCTACCGATGGTCTGGTTGACAGCGCTGGCACTGGGGACCTCTCTTACG GTTACCAGGGCCGCTCCTTTGAACCTGTAGGTACTCGGCCCCGAGTGGACTCCATGAGCT CTGTGGAGGAGGATGACTACGACACATTGACCGACATCGATTCCGACAAGAATGTCATTC GCACCAAGCAATACCTCTATGTGGCTGACCTGGCACGGAAGGACAAGCGTGTTCTGCGGA AAAAGTACCAGATCTACTTCTGGAACATTGCCACCATTGCTGTCTTCTATGCCCTTCCTG TGGTGCAGCTGGTGATCACCTACCAGACGGTGGTGAATGTCACAGGGAATCAGGACATCT GCTACTACAACTTCCTCTGCGCCCACCCACTGGGCAATCTCAGCGCCTTCAACAACATCC TCAGCAACCTGGGGTACATCCTGCTGGGGCTGCTTTTCCTGCTCATCATCCTGCAACGG AGATCAACCACAACCGGGCCCTGCTGCGCAATGACCTCTGTGCCCTGGAATGTGGGATCC CCAAACACTTTGGGCTTTTCTACGCCATGGGCACAGCCCTGATGATGGAGGGGCTGCTCA GTGCTTGCTATCATGTGTGCCCCAACTATACCAATTTCCAGTTTGACACATCGTTCATGT ACATGATCGCCGGACTCTGCATGCTGAAGCTCTACCAGAAGCGGCACCCGGACATCAACG CCAGCGCCTACAGTGCCTACGCCTGCCTGGCCATTGTCATCTTCTTCTCTGTGCTGGGCG TGGTCTTTGGCAAAGGGAACACGCGTTCTGGATCGTCTTCTCCATCATTCACATCATCG CCACCTGCTCCTCAGCACGCAGCTCTATTACATGGGCCGGTGGAAACTGGACTCGGGGA TCTTCCGCCGCATCCTCCACGTGCTCTACACAGACTGCATCCGGCAGTGCAGCGGGCCGC CCTATGGGCTTATCATGCGCCCCAATGATTTCGCTTCCTACTTGTTGGCCATTGGCATCT GCAACCTGCTCCTTTACTTCGCCTTCTACATCATCATGAAGCTCCGGAGTGGGGAGAGGA TCAAGCTCATCCCCCTGCTCTGCATCGTTTGCACCTCCGTGGTCTGGGGCTTCGCGCTCT TCTTCTTCTCCAGGGACTCAGCACCTGGCAGAAAACCCCTGCAGAGTCGAGGGAGCACA ACCGGGACTGCATCCTCGACTTCTTTGACGACCACGACATCTGGCACTTCCTCCTC CCATCGCCATGTTCGGGTCCTTCCTGGTGTTGCTGACACTGGATGACGACCTGGATACTG ${\tt TGCAGCGGGACAAGATCTATGTCTTC}{\tt TAG}{\tt CAGGAGCTGGGCCCTTCGCTTCACCTCAAGG}$ GGCCTGAGCTCCTTTGTGTCATAGACCGGTCACTCTGTCGTGCTGTGGGGATGAGTCCC AGCACCGCTGCCCAGCACTGGATGGCAGCAGGACAGCCAGGTCTAGCTTAGGCTTGGCCT GGGACAGCCATGGGGTGGCATGGAACCTTGCAGCTGCCCTCTGCCGAGGAGCAGGCCTGC TCCCCTGGAACCCCAGATGTTGGCCAAATTGCTGCTTTCTTCTCAGTGTTGGGGCCTTC

CACCCTCCCCATTTCATGCCTTGCATTTTGCCCGTCCTCCCCCACAATGCCCCAGCCT GGGACCTAAGGCCTCTTTTTCCTCCCATACTCCCACTCCAGGGCCTAGTCTGGGGCCTGA ATCTCTGTCCTGTATCAGGGCCCCAGTTCTCTTTGGGCTGTCCCTGGCTGCCATCACTGC CCATTCCAGTCAGCCAGGATGGATGGGGGTATGAGATTTTGGGGGTTGGCCAGCTGGTGC CAGACTTTTGGTGCTAAGGCCTGCAAGGGGCCTGGGGCAGTGCGTATTCTCTTCCCTCTG ACCTGTGCTCAGGGCTGGCTCTTTAGCAATGCGCTCAGCCCAATTTGAGAACCGCCTTCT GATTCAAGAGGCTGAATTCAGAGGTCACCTCTTCATCCCATCAGCTCCCAGACTGATGCC AGCACCAGGACTGGAGGGAGAAGCGCCTCACCCCTTCCCTTCTTTCCAGGCCCTTA GTCTTGCCAAACCCCAGCTGGTGGCCTTTCAGTGCCATTGACACTGCCCAAGAATGTCCA GGGGCAAAGGAGGGATGATACAGAGTTCAGCCCGTTCTGCCTCCACAGCTGTGGGCACCC CAGTGCCTACCTTAGAAAGGGGCTTCAGGAAGGGATGTGCTGTTTCCCTCTACGTGCCCA GTCCTAGCCTCGCTCTAGGACCCAGGGCTGGCTTCTAAGTTTCCGTCCAGTCTTCAGGCA AGTTCTGTGTTAGTCATGCACACACATACCTATGAAACCTTGGAGTTTACAAAGAATTGC CCCAGCTCTGGGCACCCTGGCCACCTTGGTCCTTGGATCCCCTTCGTCCCACCTGGTCCA CCCCAGATGCTGAGGATGGGGGGCCTCAGGCGGGGCCTCTGCTTTGGGGATGGGAATGTG TTTTTCTCCCAAACTTGTTTTTATAGCTCTGCTTGAAGGGCTGGGAGATGAGGTGGGTCT

FIGURE 218

MFALGLPFLVLLVASVESHLGVLGPKNVSQKDAEFERTYVDEVNSELVNIYTFNHTVTRN
RTEGURVSUNVLNKQKGAPLLFVURCKEAVVSFQVPLILRGMFQRKYLYQKVERTLCQPP
TKHESE1GPFYVDVSTLSPVNTTYQLRVSRMDDFVLRTGEGPSFNTTAAQPGYFKVEFPE
GVDSVIVKVTSNKAFPCSVISIQDVLCPVYDLDNNVAFIGMYQTMTKKAAITVQRKDFPS
NSFYVVVVVKTEDQACGGSLPFYPFAEDEPVDQGHRQKTLSVLVSQAVTSEAYVGMLFC
LGIFLSFYLLTVLLACWENWRQKKKTLLVAIDRACPESGHPRVLADSFPGSSPYEGYNYG
SFENUSGSTDGLVDSAGTGDLSYGYQGRSFEPVGTRPRVDSMSSVEEDDYDTLTDIDSDK
NVIRTKQYLYVADLARKDKRVLRKKYQIYFWNIATIAVFYALPVVQLVITTQTVVNVTGN
QDICYYNFLCAHPLGNLSAFNNILSNLGYILLGLLFLLIILQREINHNRALLRNDLCALE
CGIPKHFGLFYAMGTALMMEGLLSACYHVCPNYTNFOFDTSFMYMIAGLCMLKLYCKHHD
DINASAYSAYACLAIVIFFSVLGOVFGKGNTAFWIVFSIIHIIATLLLSTDLYYMGRWKL
DSGIFRRILHVLYTDCIRQCSGPLYVDRMVLLVMGNVINWSLAAYGLIMRPNDFASYLLA
IGIONLLYFAFYIIMKLRSGERIKLIPLLCIVCTSVVWGFALFFFFGGLSTWQKTPAES
REHNRDCILLDFFFDDIWHFLSSIAMFGSFLVLLTLDDDLDTVQRDKIYYF

Important features of the protein: Signal peptide: amino acids 1-18

Transmembrane domains: amino acids 292-317, 451-470, 501-520, 607-627, 751-770

Leucine zipper pattern:

N-glycosylation sites: amino acids 27-30, 54-57, 60-63, 123-126, 141-144, 165-168, 364-367, 476-479, 496-499, 572-575, 603-606, 699-702

FIGURE 219

AATTTTTCACCAGAGTAAACTTGAGAAACCAACTGGACCTTGAGTATTGTACATTTTGCC ${\tt TCGTGGACCCAAAGGTAGCAATCTGAAAC\underline{ATG}} {\tt AGGAGTACGATTCTACTGTTTTGTCTTC}$ TGGCTCCGGATCAGGGAACACTACCAAACCAACAGCAGTCAAATCAGGTCTTTCCTTCTT TAAGTCTGATACCATTAACACAGATGCTCACACTGGGGCCAGATCTGCATCTGTTAAATC CTGCTGCAGGAATGACACCTGGTACCCAGACCCACCCATTGACCCTGGGAGGGTTGAATG TACAACAGCAACTGCACCCACATGTGTTACCAATTTTTGTCACACAACTTGGAGCCCAGG GCACTATCCTAAGCTCAGAGGAATTGCCACAAATCTTCACGAGCCTCATCATCCATTCCT TGTTCCCGGGAGGCATCCTGCCCACCAGTCAGGCAGGGGCTAATCCAGATGTCCAGGATG GAAGCCTTCCAGCAGGAGGAGCAGGTGTAAATCCTGCCACCCAGGGAACCCCAGCAGGCC GCCTCCCAACTCCCAGTGGCACAGATGACGACTTTGCAGTGACCACCCCTGCAGGCATCC AAAGGAGCACACATGCCATCGAGGAAGCCACCACAGAATCAGCAAATGGAATTCAGTAAG $\tt CTGTTTCAAATTTTTTCAACTAAGCTGCCTCGAATTTGGTGATACATGTGAATCTTTATC$ TAATTTACCTGAAAATATTCTTGAAATTTCAGAAAATATGTTCTATGTAGAGAATCCCAA CTTTTAAAAACAATAATTCAATGGATAAATCTGTCTTTGAAATATAACATTATGCTGCCT

FIGURE 220

MRSTILLFCLLGSTRSLPQLKPALGLPPTKLAPDQGTLPNQQQSNQVFPSLSLIPLTQML TLGPDLHLLNPAAGMTPGTQTHPLTLGGLNVQQQLHPHVLPIFVTQLGAQGTILSSEELP QIFTSLIIHSLFPGGILPTSQAGANPDVQDGSLPAGGAGVNPATQGTPAGRLPTPSGTDD DFAVTTPAGIORSTHAIEBATTESANGIQ

Signal peptide: amino acids 1-16

FIGURE 221

GACTTTGCTTGAATGTTTACATTTTCTGCTCGCTGTCCTACATATCACAATATAGTGTTC ACGTTTTGTTAAAACTTTGGGGTGTCAGGAGTTGAGCTTGCTCAGCAAGCCAGCATGGCT AGGATGAGCTTTGTTATAGCAGCTTGCCAATTGGTGCTGGGCCTACTAATGACTTCATTA ACCGAGTCTTCCATACAGAATAGTGAGTGTCCACAACTTTGCGTATGTGAAATTCGTCCC TGGTTTACCCCACAGTCAACTTACAGAGAAGCCACCACTGTTGATTGCAATGACCTCCGC TTAACAAGGATTCCCAGTAACCTCTCTAGTGACACACAAGTGCTTCTCTTACAGAGCAAT AACATCGCGAAGACTGTGGATGAGCTGCAGCAGCTTTTCAACTTGACTGAACTAGATTTC TCCCAAAACAACTTTACTAACATTAAGGAGGTCGGGCTGGCAAACCTAACCCAGCTCACA ACGCTGCATTTGGAGGAAAATCAGATTACCGAGATGACTGATTACTGTCTACAAGACCTC AGCAACCTTCAAGAACTCTACATCAACCACAACCAAATTAGCACTATTTCTGCTCATGCT TTTGCAGGCTTAAAAAATCTATTAAGGCTCCACCTGAACTCCAACAAATTGAAAGTTATT GATAGTCGCTGGTTTGATTCTACACCCAACCTGGAAATTCTCATGATCGGAGAAAACCCT GTGATTGGAATTCTGGATATGAACTTCAAACCCCTCGCAAATTTGAGAAGCTTAGTTTTG GCAGGAATGTATCTCACTGATATTCCTGGAAATGCTTTGGTGGGTCTGGATAGCCTTGAG AGCCTGTCTTTTTATGATAACAAACTGGTTAAAGTCCCTCAACTTGCCCTGCAAAAAGTT CCAAATTTGAAATTCTTAGACCTCAACAAAAACCCCCATTCACAAAATCCAAGAAGGGGAC TTCAAAAATATGCTTCGGTTAAAAGAACTGGGAATCAACAATATGGGCGAGCTCGTTTCT GTCGACCGCTATGCCCTGGATAACTTGCCTGAACTCACAAAGCTGGAAGCCACCAATAAC CCTAAACTCTCTTACATCCACCGCTTGGCTTTCCGAAGTGTCCCTGCTCTGGAAAGCTTG CTGCGTGAGATCAGTATCCATAGCAATCCCCTCAGGTGTGACTGTGTGATCCACTGGATT AACTCCAACAAAACCAACATCCGCTTCATGGAGCCCCTGTCCATGTTCTGTGCCATGCCG CCCGAATATAAAGGGCACCAGGTGAAGGAAGTTTTAATCCAGGATTCGAGTGAACAGTGC CTCCCAATGATATCTCACGACAGCTTCCCAAATCGTTTAAACGTGGATATCGGCACGACG GTTTTCCTAGACTGTCGAGCCATGGCTGAGCCAGAACCTGAAATTTACTGGGTCACTCCC ATTGGAAATAAGATAACTGTGGAAACCCTTTCAGATAAATACAAGCTAAGTAGCGAAGGT ACCTTGGAAATATCTAACATACAAATTGAAGACTCAGGAAGATACACATGTGTTGCCCAG AATGTCCAAGGGGCAGACACTCGGGTGGCAACAATTAAGGTTAACGGGACCCTTCTGGAT GGTA CCCAGGTGCTAAAAATATACGTCAAGCAGACAGAATCCCATTCCATCTTAGTGTCC TGGAAAGTTAATTCCAATGTCATGACGTCAAACTTAAAATGGTCGTCTGCCACCATGAAG ATTGATAACCCTCACATAACATATACTGCCAGGGTCCCAGTCGATGTCCATGAATACAAC CTAACGCATCTGCAGCCTTCCACAGATTATGAAGTGTGTCTCACAGTGTCCAATATTCAT CAGCAGACTCAAAAGTCATGCGTAAATGTCACAACCAAAAATGCCGCCTTCGCAGTGGAC ATCTCTGATCAAGAAACCAGTACAGCCCTTGCTGCAGTAATGGGGTCTATGTTTGCCGTC ATTAGCCTTGCGTCCATTGCTGTGTACTTTGCCAAAAGATTTAAGAGAAAAAACTACCAC CACTCATTAAAAAAGTATATGCAAAAAACCTCTTCAATCCCACTAAATGAGCTGTACCCA CCACTCATTAACCTCTGGGAAGGTGACAGCGAGAAAGACAAAGATGGTTCTGCAGACACC AAGCCAACCCAGGTCGACACATCCAGAAGCTATTACATGTGG<u>TAA</u>CTCAGAGGATATTTT GCTTCTGGTAGTAAGGAGCACAAAGACGTTTTTGCTTTATTCTGCAAAAGTGAACAAGTT GAAGACTTTTGTATTTTTGACTTTGCTAGTTTGTGGCAGAGTGGAGAGGACGGGTGGATA TTTCAAATTTTTTTAGTATAGCGTATCGCAAGGGTTTGACACGGCTGCCAGCGACTCTAG GCTTCCAGTCTGTGTTTGGTTTTTATTCTTATCATTATTATGATTGTTATTATATTATTA TTTTATTTTAGTTGTTGTGCTAAACTCAATAATGCTGTTCTAACTACAGTGCTCAATAAA ΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑ

FIGURE 222

MARMSFVIAACQLVLGLLMTSLTESSIQNSECPQLCVCEIRPWFTPQSTYREATTUDCND
LRITRIPSNLSSDTQVLLLQSNNIAKTVDELQQLFNLTELDFSQNNFTNIKEVGLANLTQ
LTTLHLEERQITEMTDYCLQDLSNLQELYININQISTISAHAFAGLKNLLKLHLNSNKLK
VIDSRWFDSTPNLEILMIGENPVIGILDMNFKPLANLRSLVLAGMYLTDIPCNALVGLDS
LESLSFYDNKLVKVPQLALQKVPNLKFLDLNKNPIHKIQEGDFKNMLRLKLGINNMGEL
VSVDRYALDNLPELTKLEATNNPKLSTIHRLAFRSVPALESLMLNNNALNAIYQKTVESL
PNLREISIHSNPLRCDCVIHWINSNKTNIRFMEPLSMFCAMPPEYKGHQVKEVLIQDSSE
QCLEMISHDSFPNRLNVDIGTTVFLDCRAMAEPEPEIYWVTPIGNKITVETLSDKYKLSS
EGTLEISNIQIEDSGRYTCVAQNVQGADTRVATIKVNGTLLDGTQVLKIYVKQTESHSIL
VSWKVNSNVMTSNLKWSSATMKIDNPHITYTARVPVDVHEYNLTHLQPSTDYEVCLTVSN
IHQQTQKSCVMVTTKNAAFAVDISDQETSTALAAVMGSMFAVISLASIAVYFAKFKKN
YHHSLKKYMQKTSSIPLNELYPPLINLWEGDSEKDKDGSADTKPTQVDTSRSYYMW

Important features: Signal peptide: Amino acids 1-25

Transmembrane domain: Amino acids 508-530

N-glycosylation sites: Amino acids 69-73;96-100;106-110;117-121;385-389;517-521; 582-586:611-615

Tyrosine kinase phosphorylation site: Amino acids 573-582

N-myristoylation sites: Amino acids 16-22;224-230;464-470;637-643;698-704

FIGURE 223

CAACCATGCAAGGACAGGGCAGGAGAAGAGAACCTGCAAAGACATATTTTGTTCCAAAA TGGCATCTTACCTTTATGGAGTACTCTTTGCTGTTGGCCTCTGTGCTCCAATCTACTGTG TGTCCCCGGCCAATGCCCCCAGTGCATACCCCCGCCCTTCCTCCACAAAGAGCACCCCTG CCTCACAGGTGTATTCCCTCAACACCGACTTTGCCTTCCGCCTATACCGCAGGCTGGTTT TGGAGACCCCGAGTCAGAACATCTTCTTCTCCCCTGTGAGTGTCTCCACTTCCCTGGCCA TGCTCTCCCTTGGGGCCCACTCAGTCACCAAGACCCAGATTCTCCAGGGCCTGGGCTTCA TGACTGTTCCCAGCAAAGACCTGACCTTGAAGATGGGAAGTGCCCTCTTCGTCAAGAAGG AGCTGCAGCTGCAGGCAAATTTCTTGGGCAATGTCAAGAGGCTGTATGAAGCAGAAGTCT TTTCTACAGATTTCTCCAACCCCTCCATTGCCCAGGCGAGGATCAACAGCCATGTGAAAA AGAAGACCCAAGGGAAGGTTGTAGACATAATCCAAGGCCTTGACCTTCTGACGGCCATGG TTCTGGTGAATCACATTTTCTTTAAAGCCAAGTGGGAGAAGCCCTTTCACCTTGAATATA CAAGAAAGAACTTCCCATTCCTGGTGGGCGAGCAGGTCACTGTGCAAGTCCCCATGATGC ACCAGAAAGAGCAGTTCGCTTTTGGGGTGGATACAGAGCTGAACTGCTTTGTGCTGCAGA TGGATTACAAGGGAGATGCCGTGGCCTTCTTTGTCCTCCCTAGCAAGGGCAAGATGAGGC GGTGGATAGAGGTGTTCATCCCCAGATTTTCCATTTCTGCCTCCTACAATCTGGAAACCA TCCTCCCGAAGATGGGCATCCAAAATGCCTTTGACAAAAATGCTGATTTTTCTGGAATTG CAAAGAGAGACTCCCTGCAGGTTTCTAAAGCAACCCACAAGGCTGTGCTGGATGTCAGTG AAGAGGGCACTGAGGCCACAGCAGCTACCACCACCAAGTTCATAGTCCGATCGAAGGATG GTCCCTCTTACTTCACTGTCTCCTTCAATAGGACCTTCCTGATGATGATTACAAATAAAG CCACAGACGGTATTCTCTTTCTAGGGAAAGTGGAAAATCCCACTAAATCCATGGCCTGTTAACTGATGGCACATTGCTAATGCACAAGAAATAACAAACCACATCCCTCT TTCTGTTCTGAGGGTGCATTTGACCCCAGTGGAGCTGGATTCGCTGGCAGGGATGCCACT TCCAAGGCTCAATCACCAAACCATCAACAGGGACCCCAGTCACAAGCCAACACCCATTAA CCCCAGTCAGTGCCCTTTTCCACAAATTCTCCCCAGGTAACTAGCTTCATGGGATGTTGCT GGGTTACCATATTTCCATTCCTTGGGGCTCCCAGGAATGGAAATACGCCAACCCAGGTTA GGCACCTCTATTGCAGAATTACAATAACACATTCAATAAAACTAAAATATGAATTCAAAA AAA

FIGURE 224

MASYLYGVLFAVGLCAPIYCVSPANAPSAYPRPSSTKSTPASQVYSLNTDFAFRLYRRLV LETPSQNIFFSPVSVSTSLAMLSLGAHSVTKTQILQGLGFNLTHTPESAIHQGFQHLVHK LTVPSKDLTLKMGSALFVKKELQLQANFLGNVKRLYEAEVFSTDFSNPSIAQARINSHVK KKTQGKVVDIIQGLDLLTAMVLVNHIFFKAKMEKPFHLEYTRKNFPFLVGEQVTVQVPMM HQKEQFAFGVDTELNCFVLQMDYKGDAVAFFVLPSKGKMRQLEQALSARTLIKWSHSLQK RWIEVFIPRFSISASYNLETILPKMGIQNAFDKNADFSGIAKRDSLQVSKATHKAVLDVS EEGTEATAATTTKFIVRSKDGPSYFTVSFNRTFLMMITNKATDGILFLGKVENPTKS

Signal peptide:

FIGURE 225

GGGAGAGAGGATAAATAGCAGCGTGGCTTCCCTGGCTCCTCTCTGCATCCTTCCCGACCT ${\tt TCCCAGCAAT} \underline{{\tt ATG}} {\tt CATCTTGCACGTCTGGTCGGCTCCTGCTCCTTCTGCTACTGGG}$ GGCCCTGTCTGGATGGCCGGCCAGCGATGACCCCCATTGAGAAGGTCATTGAAGGGATCAA CCGAGGGCTGAGCAATGCAGAGAGAGAGGCGCCAGGATGGCATCAACAGTGG AATCACGCATGCCGGAAGGGAAGTGGAGAAGGTTTTCAACGGACTTAGCAACATGGGGAG CCACACCGGCAAGGAGTTGGACAAAGGCGTCCAGGGGCTCAACCACGGCATGGACAAGGT TGCCCATGAGATCAACCATGGTATTGGACAAGCAGGAAAGGAAGCAGAGAAGCTTGGCCA TGGGGTCAACAACGCTGCTGGACAGGCCGGGAAGGAAGCAGACAAAGCGGTCCAAGGGTT CCACACTGGGGTCCACCAGGCTGGGAAGGAAGCAGAGAAACTTGGCCAAGGGGTCAACCA TGCTGCTGACCAGGCTGGAAAGGAAGTGGAGAAGCTTGGCCAAGGTGCCCACCATGCTGC TGGCCAGGCCGGAAGGAGCTGCAGAATGCTCATAATGGGGTCAACCAAGCCAGCAAGGA GGCCAACCAGCTGCTGAATGGCAACCATCAAAGCGGATCTTCCAGCCATCAAGGAGGGGC CACAACCACGCCGTTAGCCTCTGGGGCCTCAGTCAACACGCCTTTCATCAACCTTCCCGC $\tt CCTGTGGAGGAGCGTCGCCAACATCATGCCC\underline{TAA}ACTGGCATCCGGCCTTGCTGGGAGAA$ $\overline{\texttt{TAATGTCGCCGTTGTCACATCAGCTGACATGACCTGGAGGGGTTGGGGGTGGGGGACAGG}}$ TTTCTGAAATCCCTGAAGGGGGTTGTACTGGGATTTGTGAATAAACTTGATACACCA

FIGURE 226

MHLARLVGSCSLLLLLGALSGWAASDDPIEKVIEGINRGLSNAEREVGKALDGINSGITH AGREVEKVFNGLSNMGSHTGKELDKGVQGLNHGMDKVAHEINHGIGQAGKEAEKLGHGVN NAAGQAGKEABKAVQGFHTGVHQAGKEAEKLGGGVNHAADQAGKEVEKLGQGAHHAAGQA GKELQNAHNGVNQASKEANQLLNGNHQSGSSSHQGGATTTPLASGASVNTPFINLPALWR SVANIMP

Important features of the protein: Signal peptide: amino acids 1-25

Homologous region to circumsporozoite (CS) repeats: amino acids 35-225

FIGURE 227

GAAGTAGAGGTGTTGTGCTGAGCGGCGCTCGGCGAACTGTGTGGACCGTCTGCTGGGACT CCGGCCCTGCGTCCGCTCAGCCCCGTGGCCCCGCGCACCTACTGCCATGGAGACGCGGCCTCGTCTCGGGGCCACCTGTTTGCTGGGCTTCAGTTTCCTGCTCCTCGTCATCTCTTGA TGGACATAATGGGCTTGGAAAGGGTTTTGGAGATCATATTCATTGGAGGACACTGGAAGA TGGGAAGAAGAAGCAGCTGCCAGTGGACTGCCCCTGATGGTGATTATTCATAAATCCTG GTGTGGAGCTTGCAAAGCTCTAAAGCCCAAATTTGCAGAATCTACGGAAATTTCAGAACT CTCCCATAATTTTGTTATGGTAAATCTTGAGGATGAAGAGGAACCCAAAGATGAAGATTT CAGCCCTGACGGGGTTATATTCCACGAATCCTTTTTCTGGATCCCAGTGGCAAGGTGCA TCCTGAAATCATCAATGAGAATGGAAACCCCAGCTACAAGTATTTTTATGTCAGTGCCGA GCAAGTTGTTCAGGGGATGAAGGAAGCTCAGGAAAGGCTGACGGGTGATGCCTTCAGAAA GAAACATCTTGAAGATGAATTGTAACATGAATGTGCCCCTTCTTTCATCAGAGTTAGTGT TCTGGAAGGAAAGCAGCAGGAAGGGAATATTGAGGAATCATCTAGAACAATTAAGCCGA CCAGGAAACCTCATTCCTACCTACACTGGAAGGAGCGCTCTCACTGTGGAAGAGTTCTGC TAACAGAAGCTGGTCTGCATGTTTGTGGATCCAGCGGAGAGTGGCAGACTTTCTTCTCCT TTTCCCTCTCACCTAAATGTCAACTTGTCATTGAATGTAAAGAATGAAACCTTCTGACAC AAAA

FIGURE 228

METRPRLGATCLLGFSFLLLVISSDGHNGLGKGFGDHIHWRTLEDGKKEAAASGLPLMVI IHKSWCGACKALKPKFAESTEISELSHNFVMVNLEDEEPKLDEDFSPDGGYIPRILFLDP SGKVHPEIINENCNPSYKYFYVSAEQVVOGMKEAQBRLTGDAFRKKHLEDEL

Signal peptide:
Amino acids 1-23

Thioredoxin family proteins Homology Block: Amino acids 58-75

FIGURE 229

CCCACGCGTCCGCCCACGCGTCCGGGTGCCACTCGCGCCCGGCCGCGCTCCGGGCTTCT CTTTTCCCTCCGACGCGCCACGCTGCCCAGACATTCCGGCTGCCGGGTCTGGAGAGCTC CCCGAACCCCTCCGCGGAGAGGAGCGAGGCGCGCCAGGGTGGCCCCCGGGGCGCCTTG GTCTCGGAGAAGCGGGGACGAGGCCGGAGGATGAGCGACTGAGGGCGACGCGGGCACTGA CGCGAGTTGGGGCCGCGACTACCGGCAGCTGACAGCGCGATGAGCGACTCCCCAGAGACC CCCTAGCCCGGTGTGCGCGCCAGGCGGAGCGCGCAGGTGGGGCTGGGCTGTTAGTGGTCC GCCCACGCGGTCGCCGGCCGGCCAGGATGGGCGCTGGCAACCCGGGCCCGCGCCCGC CGCTGCTACCCCTGCGCCCGCTGCGAGCCCGGCGTCCGGCCCGCGCCCTGCGCTCATGGA CGGCGGCTCCCGGCTGGCGGCGCGCCCCCGGGCTGTGAATGCGACTCGCCCCTCGGC CGCGCTCCCGCCCGCCCGCCGGGACGTGGTAGGGGATGCCCAGCTCCACTGCGAT GGCAGTTGGCGCGCTCTCCAGTTCCCTCCTGGTCACCTGCTGCCTGATGGTGGCTCTGTG CAGTCCGAGCATCCCGCTGGAGAAGCTGGCCCAGGCACCAGAGCAGCCGGGCCAGGAGAA GCGTGAGCACGCCACTCGGGACGCCCGGGGCGGTGAACGAGCTCGGGCGCCCGGCGAG TGAGCCGTGGAGCAAGCTGAAGCAGGCCTGGGTCTCCCAGGGCGGGGGGCGCCAAGGCCGG GGATCTGCAGGTCCGGCCCCGCGGGGACACCCCGCAGGCGGAAGCCCTGGCCGCAGCCGC CCAGGACGCGATTGGCCCGGAACTCGCGCCCACGCCCGAGCCACCCGAGGAGTACGTGTA CCCGGACTACCGTGGCAAGGGCTGCGTGGACGAGAGCGGCTTCGTGTACGCGATCGGGGA GAAGTTCGCGCCGGGCCCTCGGCCTGCCCGTGCCTGTGCACCGAGGGGGCCGCTGTG CGCGCAGCCCGAGTGCCCGAGGCTGCACCCGCGCTGCATCCACGTCGACACGAGCCAGTG TTTGGAGGAGTTCGTGGTGTCTCCATGCGAGAGGTGTCGCTGTGAAGCCAACGGTGAGGT GCTATGCACAGTGTCAGCGTGTCCCCAGACGGAGTGTGTGGACCCTGTGTACGAGCCTGA TCAGTGCTGTCCCATCTGCAAAAATGGTCCAAACTGCTTTGCAGAAACCGCGGTGATCCC TGCTGGCAGAGAAGTGAAGACTGACGAGTGCACCATATGCCACTGTACTTATGAGGAAGG CACATGGAGAATCGAGCGGCAGGCCATGTGCACGAGACATGAATGCAGGCAAATGTAGAC GCTTCCCAGAACACACACTCTGACTTTTTCTAGAACATTTTACTGATGTGAACATTCTAG ATGACTCTGGGAACTATCAGTCAAAGAAGACTTTTGATGAGGAATAATGGAAAATTGTTG GTACTTTCCTTTCTTGATAACAGTTACTACAACAGAAGGAAATGGATATATTTCAAAA AGTACACAAAAGTACACTATTATATATCAAATGTATTTCTATAATCCCTCCATTAGAGAG CTTATATAAGTGTTTTCTATAGATGCAGATTAAAAATGCTGTGTTGTCAACCGTCAAAAA

FIGURE 230

MPSSTAMAVGALSSSLLVTCCLMVALCSPSIPLEKLAQAPEQPGQEKREHATRDGPGRVN
ELGRPARDEGGSGRDWKSKSGRGLAGREPWSKLKQAWVSQGGGKKAGDLQVRPRGDTPQA
EALAAAAQDAIGPELAPTPEPPBEEYVYPDYRGKGCVDESGFVYAIGBKFAPGPFSACPCLC
TEEGPLCAQPECPRLHPRCIHVDTSQCCPQCKERKNYCEFRGKTYQTLEEFVVSPCERCR
CEANGEVLCTVSACPQTECVDPVYEPDQCCPICKNGPNCFAETAVIPAGREVKTDECTIC
HCTYEEGTWRIERQAMCTRHECRQM

Important features of the protein: Signal peptide: amino acids 1-27

Transmembrane domain: amino acids 11-30

Glycosaminoglycan attachment site: amino acids 80-83

N-myristoylation sites: amino acids 10-15, 102-107, 103-108

Cell attachment sequence: amino acids 114-117

EGF-like domain cysteine pattern signature: amino acids 176-187

FIGURE 231

GGCCGGACGCCTCCGCGTTACGGGATGAATTAACGGCGGGTTCCGCACGGAGGTTGTGAC GTCACAGGTGGGAGGTGGAACTATCAGGCTGAAAAACAGAGTGGGTACTCTCTTCTGG AAGCTGGCAACAAATGGATGATGTGATATATGCATTCCAGGGGAAGGGAAATTGTGGTGC TTCTGAACCCATGGTCAATTAACGAGGCAGTTTCTAGCTACTGCACGTACTTCATAAAGC AGGACTCTAAAAGCTTTGGAATCATGGTGTCATGGAAAGGGATTTACTTTATACTGACTC TGTTTTGGGGAAGCTTTTTTGGAAGCATTTTCATGCTGAGTCCCTTTTTACCTTTGATGT TTGTAAACCCATCTTGGTATCGCTGGATCAACAACCGCCTTGTGGCAACATGGCTCACCC TACCTGTGGCATTATTGGAGACCATGTTTGGTGTAAAAGTGATTATAACTGGGGATGCAT TTGTTCCTGGAGAAGAAGTGTCATTATCATGAACCATCGGACAAGAATGGACTGGATGT TCCTGTGGAATTGCCTGATGCGATATAGCTACCTCAGATTGGAGAAAATTTGCCTCAAAG CGAGTCTCAAAGGTGTTCCTGGATTTGGTTGGGCCATGCAGGCTGCTGCCTATATCTTCA ATATTCACGAACCACTTCAACTCCTCATATTCCCAGAAGGGACTGATCTCACAGAAAACA GCAAGTCTCGAAGTAATGCATTTGCTGAAAAAAATGGACTTCAGAAATATGAATATGTTT TACATCCAAGAACTACAGGCTTTACTTTTGTGGTAGACCGTCTAAGAGAAGGTAAGAACC TTGATGCTGTCCATGATATCACTGTGGCGTATCCTCACAACATTCCTCAATCAGAGAAGC ACCTCCTCCAAGGAGACTTTCCCAGGGAAATCCACTTTCACGTCCACCGGTATCCAATAG ACACCCTCCCACATCCAAGGAGGACCTTCAACTCTGGTGCCACAAACGGTGGGAAGAGA AAGAAGAGAGGCTGCGTTCCTTCTATCAAGGGGAGAAGAATTTTTATTTTACCGGACAGA GTGTCATTCCACCTTGCAAGTCTGAACTCAGGGTCCTTGTGGTCAAATTGCTCTCTATAC TGTATTGGACCCTGTTCAGCCCTGCAATGTGCCTACTCATATATTTGTACAGTCTTGTTA AGTGGTATTTTATAATCACCATTGTAATCTTTGTGCTGCAAGAGAGAATATTTGGTGGAC TGGAGATCATAGAACTTGCATGTTACCGACTTTTACACAAACAGCCACATTTAAATTCAA AGAAAAATGAGTAAGATTATAAGGTTTGCCATGTGAAAACCTAGAGCATATTTTGGAAAT GTTCTAAACCTTTCTAAGCTCAGATGCATTTTTGCATGACTATGTCGAATATTTCTTACT GCCATCATTATTTGTTAAAGATATTTTGCACTTAATTTTGTGGGAAAAATATTGCTACAA TTTTTTTAATCTCTGAATGTAATTTCGATACTGTGTACATAGCAGGGAGTGATCGGGGT GAAATAACTTGGGCCAGAATATTATTAAACAATCATCAGGCTTTTAAA

FIGURE 232

MHSRGREIVVLLNPWSINEAVSSYCTYFIKQDSKSFGIMVSWKGIYFILTLFWGSFFGSI
FMLSPFLPLMFVNPSWYRWINNRLVATWLTLPVALLETMFGVKVIITGDAFVPGERSVII
MNHRTRMDWMFLWNCLMRYSYLRLEKICLKASLKGVPGFGWAMQAAAYIFIHRKWKDDKS
HFEDMIDYFCDIHEPLQLLIFPEGTDLTENSKSRSNAFAEKWGLQKYEYVLHPRTTGFTF
VVDRLREGKNLDAVHDITVAYPHNIPQSEKHLLQGDFPREIHFHVHRYPIDTLPTSKEDL
QLWCHKRWEEKEERLRSFYQGEKNFYFTGQSVIPPCKSELRVLVVKLLSILYWTLFSPAM
CLLIYLYSLVKWYFIITTVIFVLQERIFGGLBIIELACYRLLHKQPHLNSKKNE

Important features of the protein: Signal peptide: amino acids 1-22

Transmembrane domains: amino acids 44-63, 90-108, 354-377

FIGURE 233

CGGCTCGAGTGCAGCTGTGGGGAGATTTCAGTGCATTGCCTCCCCTGGGTGCTCTTCATC TTGGATTTGAAAGTTGAGAGCAGCATGTTTTTGCCCACTGAAACTCATCCTGCTGCCAGTG TTACTGGATTATTCCTTGGGCCTGAATGACTTGAATGTTTCCCCGCCTGAGCTAACAGTC CATGTGGGTGATTCAGCTCTGATGGGATGTGTTTTCCAGAGCACAGAAGACAAATGTATA TTCAAGATAGACTGGACTCTGTCACCAGGAGAGCACGCCAAGGACGAATATGTGCTATAC TATTACTCCAATCTCAGTGTGCCTATTGGGCGCTTCCAGAACCGCGTACACTTGATGGGG GACATCTTATGCAATGATGGCTCTCTCCTGCTCCAAGATGTGCAAGAGGCTGACCAGGGA ACCTATATCTGTGAAATCCGCCTCAAAGGGGAGAGCCAGGTGTTCAAGAAGGCGGTGGTA ATGGGATGTGTTTTCCAGAGCACAGAAGTGAAACACGTGACCAAGGTAGAATGGATATTT TCAGGACGGCGCGCAAAGGAGGAGATTGTATTTCGTTACTACCACAAACTCAGGATGTCT GTGGAGTACTCCCAGAGCTGGGGCCACTTCCAGAATCGTGTGAACCTGGTGGGGGACATT TTCCGCAATGACGGTTCCATCATGCTTCAAGGAGTGAGGGAGTCAGATGGAGGAAACTAC ACCTGCAGTATCCACCTAGGGAACCTGGTGTTCAAGAAAACCATTGTGCTGCATGTCAGC CCGGAAGAGCCTCGAACACTGGTGACCCCGGCAGCCCTGAGGCCTCTGGTCTTGGGTGGT AATCAGTTGGTGATCATTGTGGGAATTGTCTGTGCCACAATCCTGCTGCTCCCTGTTCTG ATATTGATCGTGAAGAGACCTGTGGAAATAAGAGTTCAGTGAATTCTACAGTCTTGGTG AAGAACACGAAGAAGACTAATCCAGAGATAAAAGAAAAACCCTGCCATTTTGAAAGATGT GAAGGGGAGAAACACATTTACTCCCCAATAATTGTACGGGAGGTGATCGAGGAAGAAGAA CCAAGTGAAAAATCAGAGGCCACCTACATGACCATGCACCCAGTTTGGCCTTCTCTGAGG TCAGATCGGAACAACTCACTTGAAAAAAAGTCAGGTGGGGGAATGCCAAAAAACACAGCAA GCCTTTTGAGAAGAATGGAGAGTCCCTTCATCTCAGCAGCGGTGGAGACTCTCTCCTGTG TGTGTCCTGGGCCACTCTACCAGTGATTTCAGACTCCCGCTCTCCCAGCTGTCCTCCTGT CTCATTGTTTGGTCAATACACTGAAGATGGAGAATTTGGAGCCTGGCAGAGAGACTGGAC AGCTCTGGAGGAACAGGCCTGCTGAGGGGAGGGAGCATGGACTTGGCCTCTGGAGTGGG ACACTGGCCCTGGGAACCAGGCTGAGCTGAGTGGCCTCAAACCCCCCGTTGGATCAGACC CCCAAATCAA

FIGURE 234

MFCPLKLILLPVLLDYSLGLNDLNVSPPELTVHVGDSALMGCVFQSTEDKCIFKIDWTLS
PGEHAKDEYVLYYYSNLSVPIGRFQNRVHLMGDILCNDGSLLLQDVQEADQGTYICEIR
KGESQVFKKAVVLHVLPEPKELMVHVGGLIQMGCVFQSTEVKHVTKVEWIFSGRRAKEE
IVFRYYHKLRMSVEYSQSWGHFQNRVNLVGDIFRNDGSIMLQGVRESDGGNYTCSIHLGN
LVFKKTIVLHVSPEPRTLVTPAALRPLVLGGNQLVIIVGIVCATILLLPVLILIVKKTC
GNKSSVNSTVLVKNTKKTNPEIKEKPCHFERCEGKHIYSPIIVREVIEEEEPSEKSEAT
YMTMHPWPSLRSDRNNSLEKKSGGGMPKTOOAF

FIGURE 235

TAAAACAGCTACAATATTCCAGGGCCAGTCACTTGCCATTTCTCATAACAGCGTCAGAGA
GAAAGAACTGACTGAAACGTTTGAGATGAAGAAGAAGTTCTCCTCCTGATCACAGCCATCTT
GGCAGTGGCTGTTGGTTTCCCAGGTCTCAGACAGAGCAAGGAACAGGAAAAAAGAAGAGTATCAG
TGACAGCGATGAATTAGCTTCAGGGTTTTTTGTGTTCCCCTTACCCATATCCATTTCCACC
ACTTCCACCAATTCCATTTCCAAGATTTCCATGGTTTAAGACTAATTTCCTATTCCAAT
ACCTGAATCTGCCCCTTCCATCACCAAAAGTAAACAAGAAGAATAAGTCA
CGATAAACCTGGTCACCTGAAATTGAAATTGACCACTTCCTTGAAGAATCAAAATTCCT
GTTAATAAAAAGAAAAAATGTAATTGAAATAGCACACAGCATTCCTCATGTCAAATATCT
TTAGTGATCTTCTTTAATTATAAACATGAAAGCAAAAGATTTTTGTTATTCTTAATTTCCACA

FIGURE 236

MKKVLLLITAILAVAVGFPVSQDQEREKRSISDSDELASGFFVFPYPYPFRPLPPIPFPR FPWFRRNFPIPIPESAPTTPLPSEK

Important features of the protein: Signal peptide: amino acids 1-17

Homologous region to B3-hordein: amino acids 47-85

FIGURE 237

TCGCCATGCCTCTGCCGGAATGCAGATCCTGGGAGTCGTCCTGACACTGCTGGGCTGGG TGAATGGCCTGGTCTCCTGTGCCCTGCCCATGTGGAAGGTGACCGCTTTCATCGGCAACA GCATCGTGGTGGCCCAGGTGGTGTGGGAGGGCCTGTGGATGTCCTGCGTGGTGCAGAGCA CCGGCCAGATGCAGTGCAAGGTGTACGACTCACTGCTGGCGCTGCCACAGGACCTGCAGG CTGCACGTGCCCTCTGTGTCATCGCCCTCCTTGTGGCCCTGTTCGGCCTGCTGGTCTACC TTGCTGGGGCCAAGTGTACCACCTGTGTGGAGGAGAAGGATTCCAAGGCCCGCCTGGTGC TCACCTCTGGGATTGTCTTTTGTCATCTCAGGGGTCCTGACGCTAATCCCCGTGTGCTGCA CGGCGCATGCCATCATCCGGGACTTCTATAACCCCCTGGTGGCTGAGGCCCAAAAGCGGG GGTTGCTGTGCCACTTGCCCCTCGGGGGGGTCCCAGGGCCCAGCCATTACATGGCCC GCTACTCAACATCTGCCCCTGCCATCTCTCGGGGGCCCTCTGAGTACCCTACCAAGAATT ACGTCTGACGTGGAGGGGAATGGGGGCTCCGCTGGCGCTAGAGCCATCCAGAAGTGGCAG TGCCCAACAGCTTTGGGATGGGTTCGTACCTTTTGTTTCTGCCTCCTGCTATTTTTCTTT TGACTGAGGATATTTAAAATTCATTTGAAAACTGAGCCAAGGTGTTGACTCAGACTCTCA CTTAGGCTCTGCTGTTTCTCACCCTTGGATGATGGAGCCAAAGAGGGGGATGCTTTGAGAT TCTGGATCTTGACATGCCCATCTTAGAAGCCAGTCAAGCTATGGAACTAATGCGGAGGCT GCTTGCTGCTGCCTTTGCAACAAGACAGACTGTCCCCAAGAGTTCCTGCTGCTGCTGG GGGCTGGGCTTCCCTAGATGTCACTGGACAGCTGCCCCCCATCCTACTCAGGTCTCTGGA GCTCCTCTCTCACCCCTGGAAAAACAAATCATCTGTTAACAAAGGACTGCCCACCTCCG GAACTTCTGACCTCTGTTTCCTCCGTCCTGATAAGACGTCCACCCCCAGGGCCAGGTCC CAGCTATGTAGACCCCCGCCCCCACCTCCAACACTGCACCCTTCTGCCCTGCCCCCCTCG TCTCACCCCTTTACACTCACATTTTTATCAAATAAAGCATGTTTTGTTAGTGCA

248/392 .

FIGURE 238

MASAGMQILGVVLTLLGWVNGLVSCALPMWKVTAFIGNSIVVAQVVWEGLWMSCVVQSTG QMQCKVYDSLLALPQDLQAARALCVIALLVALFGLLVYLAGAKCTTCVEEKDSKARLVLT SGIVFVISGVLTLIPVCWTAHAIIRDFYNPLVAEAQKRELGASLYLGWAASGLLLLGGGL LCCTCPSGGSQGPSHYMARYSTSAPAISRGPSEYPYKNYV

Transmembrane domains:

amino acids 8-30 (type II), 82-102, 121-140, 166-186

FIGURE 239

FIGURE 240

MKITGGLLLLCTVVYFCSSSEAASLSPKKVDCSIYKKYPVVAIPCPITYLPVCGSDYITYGNECHLCTESLKSNGRVOFLHDGSC

Signal peptide: amino acids 1-19

FIGURE 241

FIGURE 242

MAASPARPAVLALTGLALLLLLCWGPGGISGNKLKLMLQKREAPVPTKTKVAVDENKAKE FLGSLKRQKRQLWDRTRPEVQQWYQOFLYMGFDEAKFEDDITYWLNRDRNGHEYYGDYYQ RHYDEDSATGPRSPYGFRHGASVNYDDY

Signal peptide: amino acids 1-30

FIGURE 243

CTCCATTAAACCACCACCAGCTCCCCAAGCCACCCCTTCAGCCATGAAGTTCCTGCTCCT ATGTGTGAGTAACACCCCAGGATACTGCAGGACATGTTGCCACTGGGGGGAGACAGCATT CCTACCACAGCTCATCGGTAACCACTGGCAATCAAGGAGAAGAAACACACAAAGGAAAGA CAAGAAGCAACAAACGACCGTAACATCA<u>TAA</u>TAACCACTGCTATCGCCTCCACCAACTCA GAGAAATATCATTTCCACAGTTCCAATTCCTCCTACATTGCTGAGTACTAGCCAAGGCTC CTCTTTATGGGGCAGATATCTATAGCCAACCCCAAAACTTCTGTCTTCTATCATTCTGTC ATTCATCTAGTAACTAATTTGGAGTTTGTATCTATCTTACGAGAACAATCATCATGCAGA TTCGTCCACAGGGGATCTGTCAGTTTGGGTCCTCCAAATGAAAAATGTCAAGACAGAATT GGACATGCAAAAGATTGACTGGGAGAACACACCTCTGATGGACAAAGGTGAGACAGAGCA GCCACAGGCAGGGAGAGCCTTCAGACTGCAACGCTGGCCTGATACGTGTCAAAGGAGAGA GGGATAGAGGAGGATTGAATAGAAGGAGACTAAGACTGCAGCTCTAAGAAAGTCTCAGCC AAACAGATGGGGAGGCCCAAAGCAAGGCTTGCCCCTCAGAGGAGCTCACGCAGGGCAGGA ATAGCCAGGTTCTCATATCCCAGGGGTTCAGACTTGGCTGAGAACAGCCCCTGGAGAACA TGGGGTGACTGCTACCATAGGTCTGGAAGTATGAGGCTGTCCACCAACTATCCCCTTGAA GCA AGTTCTCTTGA AGGAAATCTAAACAGTGCACCCCCATGGCTGCCACGGAGTATAAG TATTTATTTATTCATTTGAGTAACAAAGCAGACAGAATACATAGCCACCATTGGTAGTAC ACCCCAAAAGCAAGGATGGCATGATGCTGGTGACTCAAACGTGCCTACTCATGGTGTCAA ATTGGCATAATCCTCTTGGGAAGCTGTGTGGAAATAAGCACAGAAGCAGAACTCTAAT TGCTTAATCCACTAAACATTACTTCTGGGAATTGGCTCATCATAAATTATCCAAGAGAAA GCACAAAGTTATGGGCACAAAGGTTTTCCATATAATATTATTTAAAATGCTGAGAAAATG TTTATAGAATAATGGAACATAATAACATTATTCAAAATTGCATTTATGCTATAGTTGTCA AAGCAGAATTATGCATAAATTTCCTCTTACAGTTCGATGCCCATTAGTTTTATATAACAT TTATTTGACACGTACTGACTTCTATCTGAGAAGAACAAACCAAAACACTCAGGCCTAAAT AATTAAAAACGGTCCTAAAAACTAGCAAACCAGATAAGAAAAGATGTTAATGCCCATTCC CTAACTTATGTCTTAGACCAAAATTAATTCTAGATGGTTTTAAAATGACAGTGTAAAAGT AAAGTATTAAAAGATTGTGTGGTCAAATATTCAATTTAAGAGCAAGGAAATTCTTATAAA TATAACAATAGAGGCAGAACTCATGTAAGAATAAATTGATTAGGTGGTATTAAATATTAA GAGTTCATTCCTTTTTGTTTATAAATACTCTTCCGTCATATGAATAGTATTCATTTGTAT ACTGGTTTGTTGATGGACATTTGGGTTGTTCCCAGTTTATGGCTATTACAAATAAAGCTT CTATGAACATTTATGTACA

FIGURE 244

MKFLLLVLAALGFLTQVIPASAGGSKCVSNTPGYCRTCCHWGETALFMCNASRKCCISYS FLPKPDLPOLIGNHWOSRRRNTQRKDKKQQTTVTS

Important features of the protein: Signal peptide: amino acids 1-16

Transmembrane domain: amino acids 1-22

N-glycosylation site:

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 79-82

N-myristoylation site: amino acids 23-28

FIGURE 245

GGAGAGAGGCGCGCGGTGAAAGGCGCATTGATGCAGCCTGCGGCGGCCTCGGAGCGCGG CGGAGCCAGACGCTGACCACGTTCCTCTCCTCGGTCTCCTCCGCCTCCAGCTCCGCGCTG CCCGGCAGCCGGAGCCATGCGACCCCAGGGCCCCGCCGCCTCCCCGCAGCGGCTCCGCG GCCTCCTGCTGCTGCTGCTGCAGCTGCCCGCGCCGTCGAGCGCCTCTGAGATCCCCA AGGGGAAGCAAAAGGCGCAGCTCCGGCAGAGGGAGGTGGTGGACCTGTATAATGGAATGT GCTTACAAGGGCCAGCAGGAGTGCCTGGTCGAGACGGGAGCCCTGGGGCCAATGTTATTC CGGGTACACCTGGGATCCCAGGTCGGGATGGATTCAAAGGAGAAAAGGGGGAATGTCTGA GGGAAAGCTTTGAGGAGTCCTGGACACCCAACTACAAGCAGTGTTCATGGAGTTCATTGA ATTATGGCATAGATCTTGGGAAAATTGCGGAGTGTACATTTACAAAGATGCGTTCAAATA AGCGTTGGTATTTCACATTCAATGGAGCTGAATGTTCAGGACCTCTTCCCATTGAAGCTA TAATTTATTTGGACCAAGGAAGCCCTGAAATGAATTCAACAATTAATATTCATCGCACTT CTTCTGTGGAAGGACTTTGTGAAGGAATTGGTGCTGGATTAGTGGATGTTGCTATCTGGG GCATCATTATTGAAGAACTACCAAAATAAATGCTTTAATTTTCATTTGCTACCTCTTTTT TTATTATGCCTTGGAATGGTTCACTTAAATGACATTTTAAATAAGTTTATGTATACATCT GAATGAAAAGCAAAGCTAAATATGTTTACAGACCAAAGTGTGATTTCACACTGTTTTTAA AGAATACTTTCTTCATAGTCACATTCTCTCAACCTATAATTTGGAATATTGTTGTGGTCT TTTGTTTTTTCTCTTAGTATAGCATTTTTAAAAAAATATAAAAGCTACCAATCTTTGTAC

FIGURE 246

MRPQGPAASPQRLRGLLLLLLQLPAPSSASEIPKGKQKAQLRQREVVDLYNGMCLQGPA GVPGRDGSPGANVIPGTPGIPGRDGFKGEKGECLRESFEESWTPNYKQCSWSSLNYGIDL GKIAECTFTKMRSNSALRVLFSGSLRLKCRNACCQRWYFTFNGAECSGPLPIEAIIYLDQ GSPEMNSTINIHRTSSVEGLCEGIGAGLVDVAIWVGTCSDYPKGDASTGWNSVSRIIIEE LPK

Signal peptide: amino acids 1-30

Transmembrane domain: amino acids 195-217

FIGURE 247

CCCACGGCTGTCTGCACTGCCACAGCAACTTCTCCAAGAAGTTCTCCTTCTACCGCCACC ATGTGAACTTCAAGTCCTGGTGGGTGGGCGACATCCCCGTGTCAGGGGCGCTGCTCACCG ACTGGAGCGACGACGATGAAGGAGCTGCACCTGGCCATCCCCGCCAAGATCACCCGGG AGAAGCTGGACCAAGTGGCGACAGCAGTGTACCAGATGATGGATCAGCTGTACCAGGGGA AGATGTACTTCCCCGGGTATTTCCCCAACGAGCTGCGAAACATCTTCCGGGAGCAGGTGC ACCTCATCCAGAACGCCATCATCGAAAGGCACCTGGCACCAGGCAGCTGGGGAGGAGGGC AGCTCTCCAGGGAGGGACCCAGCCTAGCACCTGAAGGATCAATGCCATCACCCCGCGGG ACCTCCCTAAGTAGCCCCCAGAGGCGCTGGGAGTGTTGCCACCGCCCTCCCCTGAAGTT TGCTCCATCTCACGCTGGGGGTCAACCTGGGGACCCCTTCCCTCCGGGCCATGGACACAC ATACATGAAAACCAGGCCGCATCGACTGTCAGCACCGCTGTGGCATCTTCCAGTACGAGA CCATCTCCTGCAACAACTGCACAGACTCGCACGTCGCCTGCTTTGGCTATAACTGCGAGT AGGGCTCAGGCATCACACCCACCCGTGCCAGGGCCCTACTGTCCCTGGGGTCCCAGGCTC TTCAGGGGCGCACCACTTCCAAGCCTGTGTCCCACAGGTCCTCGGCGCAGTGGAAGTCAG AACACAGCAGTGTGTGTGTGAGCACCTCGTGGGTGAGTATGTGTGGGGCACAGGCTG GCTCCCTCAGCTCCCACGTCCTAGAGGGGCTCCCGAGGAGGTGGAACCTCAACCCAGCTC TGCGCAGGAGGCGGCTGCAGTCCTTTTCTCCCTCAAAGGTCTCCGACCCTCAGCTGGAGG CGGGCATCTTTCCTAAAGGGTCCCCATAGGGTCTGGTTCCACCCCATCCCAGGTCTGTGG TCAGAGCCTGGGAGGGTTCCCTACGATGGTTAGGGGTGCCCCATGGAGGGGCTGACTGCC CCACATTGCCTTTCAGACAGGACACGAGCATGAGGTAAGGCCGCCCTGACCTGGACTTCA GGGGGAGGGGTAAAGGGAGAGAGGGGGGGGCTAGGGGGTCCTCTAGATCAGTGGGGGC ACTGCAGGTGGGGCTCTCCCTATACCTGGGACACCTGCTGGATGTCACCTCTGCAACCAC ACCCATGTGGTGGTTTCATGAACAGACCACGCTCCTCTGCCTTCTCCTGGCCTGGGACAC ACAGAGCCACCCGGCCTTGTGAGTGACCCAGAGAAGGGAGGCCTCGGGAGAAGGGGTGC TCGTAAGCCAACACCAGCGTGCCGCGGCCTGCACACCCTTCGGACATCCCAGGCACGAGG GTGTCGTGGATGTGGCCACACATAGGACCACACGTCCCAGCTGGGAGGAGAGGCCTGGGG CCCCAGGGAGGAGGCAGGGGGTGGGGGACATGGAGAGCTGAGGCAGCCTCGTCTCCCC GCAGCCTGGTATCGCCAGCCTTAAGGTGTCTGGAGCCCCCACACTTGGCCAACCTGACCT TGGAAGATGCTGCTGAGTGTCTCAAGCAGCACTGACAGCAGCTGGGCCTGCCCCAGGGCA ACGTGGGGGGGGAGACTCAGCTGGACAGCCCCTGCCTGTCACTCTGGAGCTGGGCTGCTG CTGCCTCAGGACCCCCTCTCCGACCCCGGACAGAGCTGAGCTGGCCAGGGCCAGGAGGGC GGGAGGGAGGGAATGGGGGTGGGCTGTGCGCAGCATCAGCGCCTGGGCAGGTCCGCAGAG CTGCGGGATGTGATTAAAGTCCCTGATGTTTCTC

FIGURE 248

MALLLCLVCLTAALAHGCLHCHSNFSKKFSFYRHHVNFKSWWVGDIFVSGALLTDWSDDT MKELHLAIFAKITREKLDQVATAVYQMMDQLYQGKMYFPGYFPNELRNIFREQVHLIQNA IIERHLAPGSWGGGQLSREGPSLAPEGSMPSFRGDLP

Signal peptide: amino acids 1-15

FIGURE 249

TGGCTGCGGCGCTCTCTCGTCGCTCGCTCTCTTCTAGAGCCGAGGGACCCGG TGGCCTCGTCGCTCAGCCCCTATTTCGGCACCAAGACTCGCTACGAGGATGTCAACCCCG CCCCGCTGCACCTCGCCCTCATTCGCCACGCACCCGCTACCCCACGGTCAAACAGA TCCGCAAGCTGAGGCAGCTGCACGGGTTGCTGCAGGCCCGCGGGTCCAGGGATGGCGGGG $\tt CGGACTGGATGGACGGCAGCTAGTAGAGAAGGGACGGCAGGATATGCGACAGCTGGCGC$ TGCGTCTGGCCTCTTCCCGGCCCTTTTCAGCCGTGAGAACTACGGCCGCCTGCGGC TCATCACCAGTTCCAAGCACCGCTGCATGGATAGCAGCGCCGCCTTCCTGCAGGGGCTGT GGCAGCACTACCACCTGGCTTGCCGCCGGCCGGACGTCGCAGATATGGAGTTTGGACCTC CAACAGTTAATGATAAACTAATGAGATTTTTTGATCACTGTGAGAAGTTTTTAACTGAAG TAGAAAAAATGCTACAGCTCTTTATCACGTGGAAGCCTTCAAAACTGGACCAGAAATGC AGAACATTTTAAAAAAAGTTGCAGCTACTTTGCAAGTGCCAGTAAATGATTTAAATGCAG ATTTAATTCAAGTAGCCTTTTTCACCTGTTCATTTGACCTGGCAATTAAAGGTGTTAAAT CTCCTTGGTGTGTGTTTTTGACATAGATGATGCAAAGGTATTAGAATATTTAAATGATC TGAAACAATATTGGAAAAGAGGATATGGGTATACTATTAACAGTCGATCCAGCTGCACCT TGTTTCAGGATATCTTTCAGCACTTGGACAAAGCAGTTGAACAGAAACAAAGGTCTCAGC CAATTTCTTCTCCAGTCATCCTCCAGTTTGGTCATGCAGAGACTCTTCTTCCACTGCTTT CTCTCATGGGCTACTTCAAAGACAAGGAACCCCTAACAGCGTACAATTACAAAAAAACAAA TGCATCGGAAGTTCCGAAGTGGTCTCATTGTACCTTATGCCTCGAACCTGATATTTGTGC TTTACCACTGTGAAAATGCTAAGACTCCTAAAGAACAATTCCGAGTGCAGATGTTATTAA ATGAAAAGGTGTTACCTTTGGCTTACTCACAAGAAACTGTTTCATTTTATGAAGATCTGA AGAACCACTACAAGGACATCCTTCAGAGTTGTCAAACCAGTGAAGAATGTGAATTAGCAA GGGCTAACAGTACATCTGATGAACTATGAGTAACTGAAGAACATTTTTAATTCTTTAGGA ATCTGCAATGAGTGATTACATGCTTGTAATAGGTAGGCAATTCCTTGATTACAGGAAGCT TTTATATTACTTGAGTATTTCTGTCTTTTCACAGAAAACATTGGGTTTCTCTCTGGGTT TGGACATGAAATGTAAGAAAAGATTTTTCACTGGAGCAGCTCTCTTAAGGAGAAACAAAT CTATTTAGAGAAACAGCTGGCCCTGCAAATGTTTACAGAAATGAAATTCTTCCTACTTAT ATAAGAAATCTCACACTGAGATAGAATTGTGATTCATAATAACACTTGAAAAGTGCTGG AGATTGTTCTGCAGTTCTCTCTTTTTCCTCAGGTAGGACAGCTCTAGCATTTTCTTAA TCAGGAATATTGTGGTAAGCTGGGAGTATCACTCTGGAAGAAAGTAACATCTCCAGATGA GAATTTGAAACAAGAAACAGAGTGTTGTAAAAGGACACCTTCACTGAAGCAAGTCGGAAA GTACAATGAAAATAAATATTTTTGGTATTTATTTATGAAATATTTGAACATTTTTTTCAAT AATTCCTTTTTACTTCTAGGAAGTCTCAAAAGACCATCTTAAATTATTATATGTTTGGAC AATTAGCAACAAGTCAGATAGTTAGAATCGAAGTTTTTCAAATCCATTGCTTAGCTAACT TTTTCATTCTGTCACTTGGCTTCGATTTTTATATTTTCCTATTATATGAAATGTATCTTT TGGTTGTTTGATTTTCTTTCTTTCTTTGTAAATAGTTCTGAGTTCTGTCAAATGCCGTG

FIGURE 250

MLRAPGCLLRTSVAPAAALAAALLSSLARCSLLEPRDPVASSLSPYFGTKTRYEDVNPVL LSGPEAPWRDPELLEGTCTPVQLVALIRHGTRYPTVKQIRKLRQLHGLLQARGSRDGGAS STGSRDLGAALADWPLWYADWMDGQLVEKGRQDMRQLALRLASLFPALFSRENYGRLRLI TSSKHRCMDSSAAFLQGLWQHYHPGLPPPDVADMEFGPPTVNDKLMRFFDHCEKFLTEVE KNATALYHVEAFKTGPEMQNILKKVAATLQVPVNDLMADLIQVAFFTCSFDLAIKGVKSF WCDVPDIDDAKVLEVINDLKQYWKRGYGYTINSRSSCTLFQDIFQHLDKAVEQKQRSQPI SSPVILQFGHAETLLPLLSLMGYFKDKEPLTAYNYKKQMHRKFRSGLIVPYASNLIFVLY HCENAKTPKEQFRVQMLLNEKVLPLAYSQETVSFYEDLKNHYKDILQSCQTSEECELARA NSTSDEL

Important features: Signal sequence amino acids 1-30

N-glycosylation sites: amino acids 242-246, 481-485

N-myristoylation sites. amino acids 107-113, 113-119, 117-123, 118-124, 128-134

Endoplasmic reticulum targeting sequence: amino acids 484-489

FIGURE 251

GGAGAGCCGCGCTGGGACCGGAGTGGGGAGCGCGCGTGGAGGTGCCACCCGGCGCGGG TGGCGGAGAGATCAGAAGCCTCTTCCCCAAGCCGAGCCAACCTCAGCGGGGACCCGGGCT CAGGGACGCGGCGGCGGCGGCGACTGCAGTGGCTGGACGATGGCAGCGTCCGCCGGA GCGCTTGGGCTCTTGACAGCTGGAGTATCAGCCTTGGAAGTATATACGCCAAAAGAAATC TTCGTGGCAAATGGTACACAAGGGAAGCTGACCTGCAAGTTCAAGTCTACTACTACGACCT GGCGGGTTGACCTCAGTCTCCTGGAGCTTCCAGCCAGAGGGGGCCGACACTACTGTGTCG TTTTTCCACTACTCCCAAGGGCAAGTGTACCTTGGGAATTATCCACCATTTAAAGACAGA TTTATACACAATGGCACCTATATCTGTGATGTCAAAAACCCTCCTGACATCGTTGTCCAG CCTGGACACATTAGGCTCTATGTCGTAGAAAAAGAGAAATTTGCCTGTGTTTCCAGTTTGG GTAGTGGTGGGCATAGTTACTGCTGTGGTCCTAGGTCTCACTCTGCTCATCAGCATGATT CTGGCTGTCCTCTATAGAAGGAAAAACTCTAAACGGGATTACACTGGCTGCAGTACATCA GAGAGTTTGTCACCAGTTAAGCAGGCTCCTCGGAAGTCCCCCTCCGACACTGAGGGTCTT GTAAAGAGTCTGCCTTCTGGATCTCACCAGGGCCCAGTCATATATGCACAGTTAGACCAC TCCGGCGGACATCACAGTGACAAGATTAACAAGTCAGAGTCTGTGGTGTATGCGGATATC CGAAAGAATTAAGAGAATACCTAGAACATATCCTCAGCAAGAAACAAAACCAAACTGGAC TCTCGTGCAGAAAATGTAGCCCATTACCACATGTAGCCTTGGAGACCCAGGCAAGGACAA GTACACGTGTACTCACAGAGGGAGAGAAAGATGTGTACAAAGGATATGTATAAATATTCT ATTTAGTCATCCTGATATGAGGAGCCAGTGTTGCATGATGAAAAGATGGTATGATTCTAC ATATGTACCCATTGTCTTGCTGTTTTTTGTACTTTTTTCAGGTCATTTACAATTGGGAG ATTTCAGAAACATTCCTTTCACCATCATTTAGAAATGGTTTGCCTTAATGGAGACAATAG CAGATCCTGTAGTATTTCCAGTAGACATGGCCTTTTAATCTAAGGGCTTAAGACTGATTA GTCTTAGCATTTACTGTAGTTGGAGGATGGAGATGCTATGATGGAAGCATACCCAGGGTG GCCTTTAGCACAGTATCAGTACCATTTATTTGTCTGCCGCTTTTAAAAAATACCCATTGG CTATGCCACTTGAAAACAATTTGAGAAGTTTTTTTGAAGTTTTTTCTCACTAAAATATGGG GCAATTGTTAGCCTTACATGTTGTGTAGACTTACTTTAAGTTTGCACCCTTGAAATGTGT CATATCAATTTCTGGATTCATAATAGCAAGATTAGCAAAGGATAAATGCCGAAGGTCACT TCATTCTGGACACAGTTGGATCAATACTGATTAAGTAGAAAATCCAAGCTTTGCTTGAGA ACTTTTGTAACGTGGAGAGTAAAAAGTATCGGTTTTA

FIGURE 252

MAASAGAGAVIAAPDSRRWLWSVLAAALGLLTAGVSALEVYTPKEIFVANGTQGKLTCKF KSTSTTGGLTSVSWSFQPEGADTTVSFFHYSQGQVYLGNYPPFKDRISWAGDLDKKDASI NIEBMQFIHNGTYICDVKNPPDIVVQPGHIRLYVVEKENLPVFPVWVVVGIVTAVVLGLT LLISMILAVLYRRKNSKRDYTGCSTSESLSPVKQAPRKSPSDTEGLVKSLPSGSHQGPVI YAOLDHSGGHHSDKINKSESVVYADIRKN

Signal peptide: amino acids 1-37

Transmembrane domain: amino acids 161-183

FIGURE 253

GTGA CA CTATAGA AGAG CTATGA CGTCGCATGCA CGCGTACGTAAGCT CGGAATT CGGCT CGAGGCTGGTGGGAAGAAGCCGAGATGGCGGCAGCCAGCGCTGGGGCAACCCGGCTGCTC CTGCTCTTGCTGATGGCGGTAGCAGCGCCCAGTCGAGCCCGGGCAGCGGCTGCCGGGCC GGGACTGGTGCGCGAGGGGCTGGGGCGGAAGGTCGAGAGGGCGAGGCCTGTGGCACGGTG GGCTGCTGCTGGAGCACTCATTTGAGATCGATGACAGTGCCAACTTCCGGAAGCGGGGC TCACTGCTCTGGAACCAGCAGGATGGTACCTTGTCCCTGTCACAGCGGCAGCTCAGCGAG GAGGAGCGGGCCGACTCCGGGATGTGGCAGCCCTGAATGGCCTGTACCGGGTCCGGATC CCAAGCCGACCCGGGCCCTGGATGGCCTGGAAGCTGGTGGCTATGTCTCCTCCTTTGTC CCTGCGTGCTCCCTGGTGGAGTCGCACCTGTCGGACCAGCTGACCCTGCACGTGGATGTG GCCGGCAACGTGGTGGCGTGTCGGTGGTGACGCACCCCGGGGGCTGCCGGGGCCATGAG GTGGAGGACGTGGACCTGGAGCTGTTCAACACCTCGGTGCAGCTGCAGCCGCCCACCACA GCCCAGGCCTGAGACGGCGGCTTCATTGAGCGCCTGGAGATGGAACAGGCCCAGAAG GCCAAGAACCCCCAGGAGCAGAAGTCCTTCTTCGCCAAATACTGGATGTACATCATTCCC GTCGTCCTGTTCCTCATGATGTCAGGAGCGCCAGACACCGGGGGCCAGGGTGGGGGTGGG TTAAAAACATCGACGATACATTGAAATGTGTGAACGTTTTGAAAAGCTACAGCTTCCAGC ACTGCTCACTTGATACGTTATTCAGAAACCCAAGGAATGGCTGTCCCCATCCTCATGTGG CTGTGTGGAGCTCAGCTGTGTTGTGTGGCAGTTTATTAAACTGTCCCCCAGATCGACACG CAAAAAAAAA

FIGURE 254

MAAASAGATRLLLLLLMAVAAPSRARGSGCRAGTGARGAGAEGREGEACGTVGLLLEHSF EIDDSANFRKRGSLLWNQQDGTLSLSQRQLSEEERGRLRDVAALNGLYRVRIPRRFGALD GLEAGGYVSSFVPACSLVESHLSDQLTLHVDVAGNVVGVSVVTHPGGCRGHEVEDVDLEL FNTSVQLQPPTTAPGPETAAFIERLEMEQAQKAKNPQEQKSFFAKYWMYIIPVVLFLMMS GAPDTGGOGGGGGGGGGGGGCLCCVPPSL

Signal peptide: amino acids 1-24

Transmembrane domain: amino acids 226-243

FIGURE 255

GCGACGCGCGGCGGGCGAGAGGAAACGCGGCCCGGGCCGGGCCCGGCCCTGGAGA TGGTCCCCGGCGCGCGGGGTGTTGTCTCGTGCTCTGGCTCCCCGCGTGCGTCGCGG CCCACGGCTTCCGTATCCATGATTATTTGTACTTTCAAGTGCTGAGTCCTGGGGACATTC GATACATCTTCACAGCCACACCTGCCAAGGACTTTGGTGGTATCTTTCACACAAGGTATG AGCAGATTCACCTTGTCCCCGCTGAACCTCCAGAGGCCTGCGGGGAACTCAGCAACGGTT CTCGGGTGGTCCAGGAGCACGGCGGGCGGGCGGTGATCATCTCTGACAACGCAGTTGACA ATGACAGCTTCTACGTGGAGATGATCCAGGACAGTACCCAGCGCACAGCTGACATCCCCG CCCTCTTCCTGCTCGGCCGAGACGGCTACATGATCCGCCGCTCTCTGGAACAGCATGGGC TGCCATGGGCCATCATTTCCATCCCAGTCAATGTCACCAGCATCCCCACCTTTGAGCTGC TGCAACCGCCCTGGACCTTCTGGTAGAAGAGTTTGTCCCACATTCCAGCCATAAGTGACT CTGAGCTGGGAAGGGGAAACCCAGGAATTTTGCTACTTGGAATTTGGAGATAGCATCTGG GGACAAGTGGAGCCAGGTAGAGGAAAAGGGTTTGGGCGTTGCTAGGCTGAAAGGGAAGCC ACACCACTGGCCTTCCCCAGGGCCCCCAAGGGTGTCTCATGCTACAAGAAGAGGC AAGAGACAGGCCCCAGGGCTTCTGGCTAGAACCCGAAACAAAAGGAGCTGAAGGCAGGTG GCCTGAGAGCCATCTGTGACCTGTCACACTCACCTGGCTCCAGCCTCCCCTACCCAGGGT $\tt CTCTGCACAGTGACCTTCACAGCAGTTGTTGGAGTGGTTTAAAGAGCTGGTGTTTGGGGA$ ΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑ

FIGURE 256

MVPGAAGWCCLVLWLPACVAAHGFRIHDYLYFQVLSPGDIRYIFTATPAKDFGGIFHTRY EQIHLVPAEPPEACGELSNGFFIQDQIALVERGGCSFLSKTRVVQEHGGRAVIISDNAVD NDSFYVEMIQDSTQRTADIPALFLLGRDGYMIRRSLEQHGLPWAIISIPVNVTSIPTFEL LQPPWTFW

Signal peptide: amino acids 1-20

FIGURE 257

FIGURE 258

MHRPEAMLLLLTLALLGGPTWAGKMYGPGGGKYFSTTEDYDHEITGLRVSVGLLLVKSVQ VKLGDSWDVKLGALGGNTQEVTLQPGEYITKVFVAFQAFLRGMVMYTSKDRYFYFGKLDG QISSAYPSQEGQVLVGIYGQYQLLGIKSIGFEWNYPLEEPTTEPPVNLTYSANSPVGR

Signal peptide: amino acids 1-22

FIGURE 259

FIGURE 260

MAQSLALSLLILVLAFGIPRTQGSDGGAQDCCLKYSQRKIPAKVVRSYRKQEPSLGCSIP AILFLPRKRSQAELCADPKELWVQQLMQHLDKTPSPQKPAQGCRKDRGASKTGKKGKGSK GCKRTERSOTPKGP

Important features of the protein: Signal peptide: amino acids 1-17

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 67-71

N-myristoylation sites: amino acids 17-23, 23-29, 27-33, 108-114, 118-124, 121-127

Amidation site: amino acids 112-116

Small cytokines: amino acids 51-91

FIGURE 261

GGGACTACAAGCCGCGCGCGCTGCCGCTGGCCCTCAGCAACCCTCGACATGGCGCTGA GGCGGCCACCGCGACTCCGGCTCTGCCTCGCTGACTTCTTCCTGCTGCTTCTTT TCAGGGGCTGCCTGATAGGGGCTGTAAATCTCAAATCCAGCAATCGAACCCCAGTGGTAC AGGAATTTGAAAGTGTGGAACTGTCTTGCATCATTACGGATTCGCAGACAAGTGACCCCA GGATCGAGTGGAAGAAATTCAAGATGAACAAACCACATATGTGTTTTTTGACAACAAAA TTCAGGGAGACTTGGCGGGTCGTGCAGAAATACTGGGGAAGACATCCCTGAAGATCTGGA ATGTGACACGGAGAGACTCAGCCCTTTATCGCTGTGAGGTCGTTGCTCGAAATGACCGCA AGGA A TTGATGAGATTGTGATCGAGTTAACTGTGCAAGTGAAGCCAGTGACCCCTGTCT GTAGAGTGCCGAAGGCTGTACCAGTAGGCAAGATGGCAACACTGCACTGCCAGGAGAGTG AGGGCCACCCCGGCCTCACTACAGCTGGTATCGCAATGATGTACCACTGCCCACGGATT CCAGAGCCAATCCCAGATTTCGCAATTCTTCTTTCCACTTAAACTCTGAAACAGGCACTT TGGTGTTCACTGCTGTTCACAAGGACGACTCTGGGCAGTACTACTGCATTGCTTCCAATG ACGCAGGCTCAGCCAGGTGTGAGGAGCAGGAGATGGAAGTCTATGACCTGAACATTGGCG GAATTATTGGGGGGGTTCTGGTTGTCCTTGCTGTACTGGCCCTGATCACGTTGGGCATCT GCTGTGCATACAGACGTGGCTACTTCATCAACAATAAACAGGATGGAGAAAGTTACAAGA ACCCAGGGAAACCAGATGGAGTTAACTACATCCGCACTGACGAGGAGGGCGACTTCAGAC ACA ACTCATCCTTTGTGATCTGAGACCCGCGGTGTGGCTGAGAGCGCACAGAGCGCACGT GCACATACCTCTGCTAGAAACTCCTGTCAAGGCAGCGAGAGCTGATGCACTCGGACAGAG CTAGACACTCATTCAGAAGCTTTTCGTTTTGGCCAAAGTTGACCACTACTCTTCTTACTC TAACAAGCCACATGAATAGAAGAATTTTCCTCAAGATGGACCCGGTAAATATAACCACAA GGAAGCGAAACTGGGTGCGTTCACTGAGTTGGGTTCCTAATCTGTTTCTGGCCTGATTCC CGCATGAGTATTAGGGTGATCTTAAAGAGTTTGCTCACGTAAACGCCCGTGCTGGGCCCT GTGAAGCCAGCATGTTCACCACTGGTCGTTCAGCAGCCACGACAGCACCATGTGAGATGG CGAGGTGGCTGGACAGCACCAGCAGCGCATCCCGGCGGGAACCCAGAAAAGGCTTCTTAC ACAGCAGCCTTACTTCATCGGCCCACAGACACCACCGCAGTTTCTTCTTAAAGGCTCTGC TGATCGGTGTTGCAGTGTCCATTGTGGAGAAGCTTTTTGGATCAGCATTTTGTAAAAACA ACCAAAATCAGGAAGGTAAATTGGTTGCTGGAAGAGGGATCTTGCCTGAGGAACCCTGCT ${\tt TGTCCAACAGGGTGTCAGGATTTAAGGAAAACCTTCGTCTTAGGCTAAGTCTGAAATGGT}$ ${ t ACTGAAATATGCTTTCTATGGGTCTTGTTTATTTATAAAATTTTACATCTAAATTTTT$ GCTAAGGATGTATTTTGATTATTGAAAAGAAAATTTCTATTTAAACTGTAAATATATTGT CATACAATGTTAAATAACCTATTTTTTAAAAAAGTTCAACTTAAGGTAGAAGTTCCAAG CTACTAGTGTTAAATTGGAAAATATCAATAATTAAGAGTATTTTACCCAAGGAATCCTCT CATGGAAGTTTACTGTGATGTTCCTTTTCTCACACAAGTTTTAGCCTTTTTCACAAGGGA ACTCATACTGTCTACACATCAGACCATAGTTGCTTAGGAAACCTTTAAAAATTCCAGTTA AGCAATGTTGAAATCAGTTTGCATCTCTTCAAAAGAAACCTCTCAGGTTAGCTTTGAACT GCCTCTTCCTGAGATGACTAGGACAGTCTGTACCCAGAGGCCACCCAGAAGCCCTCAGAT GTACATACACAGATGCCAGTCAGCTCCTGGGGTTGCGCCAGGCGCCCCCGCTCTAGCTCA CTGTTGCCTCGCTGTCTGCCAGGAGGCCCTGCCATCCTTGGGCCCTGGCAGTGGCTGTGT CCCAGTGAGCTTTACTCACGTGGCCCTTGCTTCATCCAGCACAGCTCTCAGGTGGGCACT GCAGGGACACTGGTGTCTTCCATGTAGCGTCCCAGCTTTGGGCTCCTGTAACAGACCTCT GTTTAATTATTTGTTAAGATTGTCTAAGGCCAAAGGCAATTGCGAAATCAAGTCTGTCAA GTACAATAACATTTTTAAAAGAAAATGGATCCCACTGTTCCTCTTTGCCACAGAGAAAGC ACCCAGACGCCACAGGCTCTGTCGCATTTCAAAACAAACCATGATGGAGTGGCGGCCAGT CCAGCCTTTTAAAGAACGTCAGGTGGAGCAGCCAGGTGAAAGGCCTGGCGGGGAGGAAAG TGAAACGCCTGAATCAAAAGCAGTTTTCTAATTTTGACTTTAAATTTTTCATCCGCCGGA

FIGURE 262

MALRRPPRIRLCARLPDFFLILLFRGCLIGAVNLKSSNRTPVVQEFESVELSCIITDSQT SDPRIEWKKIQDEQTTYVFFDNKIQGDLAGRAEILGKTSLKIWNVTRRDSALYRCEVVAR NDRKEIDEIVIELTVQVKPVTPVCRVPKAVPVGKMATLHCQESEGHPRPHYSWYRNDVPL PTDSRANPRFRNSSFHLNSETGTLVFTAVHKDDSGQYYCIASNDAGSARCEEQEMEVYDL NIGGIIGGVLVVLAVLALITLGICCAYRRGYFINNKQDGESYKNPGKPDGVNYIRTDEEG DFRIKSSFVI

Important features of the protein: Signal peptide: amino acids 1-30

Transmembrane domain: amino acids 243-263

N-glycosylation sites: amino acids 104-107, 192-195

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 107-110

Casein kinase II phosphorylation site: amino acids 106-109, 296-299

Tyrosine kinase phosphorylation site: amino acids 69-77

N-myristoylation sites: amino acids 26-31, 215-220, 226-231, 243-248, 244-249, 262-267

FIGURE 263

CCAGGACCAGGGCGCACCGGCTCAGCCTCTCACTTGTCAGAGGCCGGGGAAGAGAAGCAA AGCGCAACGGTGTGGTCCAAGCCGGGGCTTCTGCTTCGCCTCTAGGACATACACGGGACC CCCTAACTTCAGTCCCCCAAACGCGCACCCTCGAAGTCTTGAACTCCAGCCCCGCACATC CACGCGCGCACAGGCGCGCAGGCGGCAGGTCCCGGCCGAAGGCGATGCGCGCAGGGGG TCGGGCAGCTGGGCTCGGGCGGGGGGGTAGGGCCCGGCAGGGAGGCAGGGAGGCTGCAT ATTCAGAGTCGCGGGCTGCGCCCTGGGCAGAGGCCGCCCTCGCTCCACGCAACACCTGCT GCTGCCACCGCGCGCGATGAGCCGCGTGGTCTCGCTGCTGCTGGGCGCCCGCGCTGCTCT GCGGCCACGGAGCCTTCTGCCGCCGCGTGGTCAGCGGCCAAAAGGTGTGTTTTGCTGACT TCAAGCATCCCTGCTACAAAATGGCCTACTTCCATGAACTGTCCAGCCGAGTGAGCTTTC AGGAGGCACGCCTGGCTTGTGAGAGTGAGGGAGGAGTCCTCCTCAGCCTTGAGAATGAAG CAGAACAGAAGTTAATAGAGAGCATGTTGCAAAACCTGACAAAACCCGGGACAGGGATTT CTGATGGTGATTTCTGGATAGGGCTTTGGAGGAATGGAGATGGGCAAACATCTGGTGCCT GCCCAGATCTCTACCAGTGGTCTGATGGAAGCAATTCCCAGTACCGAAACTGGTACACAG CTGGCCTTGGGGGTCCCTACCTTTACCAGTGGAATGATGACAGGTGTAACATGAAGCACA ATTATATTTGCAAGTATGAACCAGAGATTAATCCAACAGCCCCTGTAGAAAAGCCTTATC TTACAAATCAACCAGGAGACACCCATCAGAATGTGGTTGTTACTGAAGCAGGTATAATTC CCAATCTAATTTATGTTGTTATACCAACAATACCCCTGCTCTTACTGATACTGGTTGCTT TTGGAACCTGTTGTTTCCAGATGCTGCATAAAAGTAAAGGAAGAACAAAAACTAGTCCAA ${\tt AACTCATTGACTTGGTTCCAGAATTTTGTAATTCTGGATCTGTATAAGGAATGGCATCAG}$ AACAATAGCTTGGAATGGCTTGAAATCACAAAGGATCTGCAAGATGAACTGTAAGCTCCC CCTTGAGGCAAATATTAAAGTAATTTTTATATGTCTATTATTTCATTTAAAGAATATGCT GTGCTAATAATGGAGTGAGACATGCTTATTTTGCTAAAGGATGCACCCAAACTTCAAACT TCAAGCAAATGAAATGGACAATGCAGATAAAGTTGTTATCAACACGTCGGGAGTATGTGT GTTAGAAGCAATTCCTTTATTTCTTTCACCTTTCATAAGTTGTTATCTAGTCAATGTAA TGTATATTGTATTGAAATTTACAGTGTGCAAAAGTATTTTACCTTTGCATAAGTGTTTGA TAAAAATGAACTGTTCTAATATTTATTTTTATGGCATCTCATTTTTCAATACATGCTCTT ATAGAAAAAGTTTGTTTTCTCGAAATAATTCATCTTTCAGCTTCTCTGCTTTTGGTCAAT GTCTAGGAAATCTCTTCAGAAATAAGAAGCTATTTCATTAAGTGTGATATAAACCTCCTC AAACATTTTACTTAGAGGCAAGGATTGTCTAATTTCAATTGTGCAAGACATGTGCCTTAT AATTATTTTTAGCTTAAAATTAAACAGATTTTGTAATAATGTAACTTTGTTAATAGGTGC ATAAACACTAATGCAGTCAATTTGAACAAAAGAAGTGACATACACAATATAAATCATATG TCTTCACACGTTGCCTATATAATGAGAAGCAGCTCTCTGAGGGTTCTGAAATCAATGTGG TCCCTCTCTTGCCCACTAAACAAGATGGTTGTTCGGGGTTTGGGATTGACACTGGAGGC AGATAGTTGCAAAGTTAGTCTAAGGTTTCCCTAGCTGTATTTAGCCTCTGACTATATTAG TATACAAAGAGGTCATGTGGTTGAGACCAGGTGAATAGTCACTATCAGTGTGGAGACAAG CACAGCACACAGACATTTTAGGAAGGAAAGGAACTACGAAATCGTGTGAAAATGGGTTGG AACCCATCAGTGATCGCATATTCATTGATGAGGGTTTGCTTGAGATAGAAAATGGTGGCT CCTTTCTGTCTTATCTCCTAGTTTCTTCAATGCTTACGCCTTGTTCTTCTCAAGAGAAAG TTGTAACTCTCTGGTCTTCATATGTCCCTGTGCTCCTTTTAACCAAATAAAGAGTTCTTG

FIGURE 264

MSRVVSLLLGAALLCGHGAFCRRVVSGQKVCFADFKHPCYKMAYFHELSSRVSFQEARLA CESEGGVLLSLENEAEQKLIESMLQNLTKPGTGISDGDFWIGLWRNGDGQTSGACPDLYQ WSDGSNSQYRNWYTDEPSCGSEKCVVWYHQPTANPGLGGPYLYQWNDDRCMMKHNYICKY EPEINPTAPVEKPYLTNQPGDTHQNVVVTEAGIIPNLIYVVIPTIPLLLLILVAFGTCCF OMLHKSKGFTKTSPNOSTLWISKSTRKESGMEV

Important features of the protein: Signal peptide: amino acids 1-21

Transmembrane domain: amino acids 214-235

N-glycosylation sites: amino acids 86-89 and 255-258

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 266-269

N-myristoylation sites: amino acids 27-32, 66-71, 91-96, 93-98, 102-107, 109-114, 140-145 and 212-217

FIGURE 265

TCCTGGCATGCCTGCCACAGCCACTGGGCCCGAAGTTGCTCAGCCTGAAGTAGACACCA CCCTGGGTCGTGTGCGAGGCCGGCAGGTGGGCGTGAAGGGCACAGACCGCCTTGTGAATG TCTTTCTGGGCATTCCATTTGCCCAGCCGCCACTGGGCCCTGACCGGTTCTCAGCCCCAC ACCCAGCACAGCCCTGGGAGGGTGTGCGGGATGCCAGCACTGCGCCCCCAATGTGCCTAC AAGACGTGGAGAGCATGAACAGCAGCAGATTTGTCCTCAACGGAAAACAGCAGATCTTCT CCGTTTCAGAGGACTGCCTGGTCCTCAACGTCTATAGCCCAGCTGAGGTCCCCGCAGGGT CCGGTAGGCCGGTCATGGTATGGGTCCATGGAGGCGCTCTGATAACTGGCGCTGCCACCT CCTACGATGGATCAGCTCTGGCTGCCTATGGGGATGTGGTCGTGGTTACAGTCCAGTACC GCCTTGGGTCCTTGGCTTCTTCAGCACTGGAGATGAGCATGCACCTGGCAACCAGGGCT TCCTAGATGTGGTAGCTGCTTTGCGCTGGGTGCAAGAAACATCGCCCCCTTCGGGGGTG ACCTCAACTGTCTCTCTTTGGTGGATCTGCCGGTGGGAGCATCATCTCTGGCCTGG TCCTGTCCCCAGTGGCTGCAGGGCTGTTCCACAGAGCCATCACACAGAGTGGGGTCATCA CCACCCCAGGGATCATCGACTCTCACCCTTGGCCCCTAGCTCAGAAAATCGCAAACACCT CTGTCTTCCCCAAAAGCCCCAAGGAACTCCTGAAGGAGAAGCCCTTCCACTCTGTGCCCT ${\tt TCCTCATGGGTGTCAACAACCATGAGTTCAGCTGGCTCATCCCCAGGGGGTTGGGGTCTCC}$ TGGATACAATGGAGCAGATGAGCCGGGAGGACATGCTGGCCATCTCAACACCCGTCTTGA CCAGTCTGGATGTGCCCCCTGAGATGATGCCCACCGTCATAGATGAATACCTAGGAAGCA ACTCGGACGCACAAGCCAAATGCCAGGCGTTCCAGGAATTCATGGGTGACGTATTCATCA ATGTTCCCACCGTCAGTTTTTCAAGATACCTTCGAGATTCTGGAAGCCCTGTCTTTTTCT ATCATGGGGCCGAGGGTGCTTTTGTGTTCCGAGGTCCCTTCCTCATGGACGAGAGCTCCCC GCCTGGCCTTTCCAGAGGCCACAGAGGAGAAGCAGCTAAGCCTCACCATGATGGCCC AGTGGACCCACTTTGCCCGGACAGGGGACCCCAATAGCAAGGCTCTGCCTCCTTGGCCCC AATTCAACCAGGCGGAACAATATCTGGAGATCAACCCAGTGCCACGGGCCGGACAGAAGT TCAGGGAGGCCTGGATGCAGTTCTGGTCAGAGACGCTCCCCAGCAAGATACAACAGTGGC ACCAGAAGCAGAAGAACAGGAAGGCCCAGGAGGACCTCTGAGGCCAGGCCTGAACCTTCT TGGCTGGGGCAAACCACTCTTCAAGTGGTGGCAGAGTCCCAGCACGGCAGCCCGCCTCTC CCCCTGCTGAGACTTTAATCTCCACCAGCCCTTAAAGTGTCGGCCGCTCTGTGACTGGAG TTATGCTCTTTTGAAATGTCACAAGGCCGCCTCCCACCTCTGGGGCATTGTACAAGTTCT TCCCTCTCCCTGAAGTGCCTTTCCTGCTTTCTTCGTGGTAGGTTCTAGCACATTCCTCTA GCTTCCTGGAGGACTCACTCCCCAGGAAGCCTTCCCTGCCTTCTCTGGGCTGTGCGGCCC CCCCCTCAGAGGAGCTCTCTCAAAATGGGGATTAGCCTAACCCCACTCTGTCACCCACAC ATATGGGAATGGCAGCTGCTGAACTTGAACCCAGAGCCTTCAGGTGCCAAAGCCATACTC AGGCCCCACCGACATTGTCCACCCTGGCCAGAAGGGTGCATGCCAATGGCAGAGACCTG GGATGGGAGAGTCCTGGGGCGCCAGGGGATCCAGCCTAGAGCAGACCTTAGCCCCTGAC GCACAAGACAACAGAATCCATCAGGGCCATGAGTGTCACCCAGACCTGACCCTCACCAAT ${\tt TCCAGCCCTGACCCTCAGGACGCTGGATGCCAGCTCCCAGCCCCAGTGCCGGGTCCTCC}$ ACAGCAGGACAGGCCAGGGGAGGGCATCTGGACCAGGGCATCCGTCGGGCTATTGTCACA GAGAAAAGAAGACCCACCCACTCGGGCTGCAAAAGGTGAAAAGCACCAAGAGGTTTTC

AGATGGAAGTGAGAGGTGACAGTGTGCTGGCAGCCCTCACAGCCCTCGCTTGCTCTCCCT GCCGCCTCTGCCTGGGCTCCCACTTTGGCAGCACTTGAGGAGCCCTTCAACCCGCCGCTG CACTGTAGGAGCCCCTTTCTGGGCTGGCCAAGGCCGGAGCCAGCTCCCTCAGCTTGCGGG GAGGTGCGGAGGGAGAGGGGCGGGCAGGAACCGGGGCTGCGCGCAGCGCTTGCGGGCCAG AGGCAGTGAGGGCCTTAGCACCTGGGCCAGCAGCTGCTGTGCTCGATTTCTCGCTGGGCC TTAGCTGCCTCCCGCGGGGCAGGGCTCGGGACCTGCAGCCCTCCATGCCTGACCCTCCC CCCACCCCGTGGGCTCCTGTGCGGCCGGAGCCTCCCCAAGGAGCGCCGCCCCTGCTC CACAGCGCCCAGTCCCATCGACCACCCAAGGGCTGAGGAGTGCGGGTGCACAGCGCGGGA CTGGCAGGCAGCTCCACCTGCTGCCCCAGTGCTGGATCCACTGGGTGAAGCCAGCTGGGC TCCTGAGTCTGGTGGGGACTTGGAGAACCTTTATGTCTAGCTAAGGGATTGTAAATACAC CGATGGGCACTCTGTATCTAGCTCAAGGTTTGTAAACACACCAATCAGCACCCTGTGTCT AGCTCAGTGTTTGTGAATGCACCAATCCACACTCTGTATCTGGCTACTCTGGTGGGGACT TGGAGAACCTTTGTGTCCACACTCTGTATCTAGCTAATCTAGTGGGGATGTGGAGAACCT TTGTGTCTAGCTCAGGGATCGTAAACGCACCAATCAGCACCCTGTCAAAACAGACCACTT GCAGGCTGCCTGAGCCAGCAGTGACAACCCCCCTCGGGTCCCCTCCCACGCCGTGGAAGC TTTGTTCTTTCGCTCTTTGCAATAAATCTTGCTACTGCCCAAAA

FIGURE 266

MERAVRVESGVLVGVVCLLLACPATATGPEVAQPEVDTTLGRVRGRQVGVKGTDRLVNVF
LGIPFAQPPLGPDRFSAPHPAQPWEGVRDASTAPPMCLQDVESMNSSRFVLNGKQQIFSV
SEDCLVLNVYSPAEVPAGSGRPVMVWVHGGALITGAATSYDGSALAAYGDVVVVTVQYRL
GVLGFFSTGDEHAPGNQGFLDVVAALRWVQENIAPFGGDLNCVTVFGGSAGGSIISGLVL
SPVAAGLFHRAITQSGVITTPGIIDSHPMPLAQKIANTLACSSSSPAEMVQCLQQKEGEE
LVLSKKLKNTIYPLTVDGTVFPKSPKELLKEKPFHSVPFLMGVNNHEFSWLIFRGWGLLD
TMEQMSREDMLAISTPVLTSLDVPPEMMPTVIDEYLGSNSDAQAKCQAFQEFMGDVFINV
PTVSFSRYLRDSGSPVFFYEFQHRPSSFAKIKPAWVKADHGAEGAFVFGGPFLMDESSRL
AFPEATEEEKQLSLTMMAQWTHFARTGDPNSKALPPWPQFNQAEQYLEINPVPRAGQKFR
EAMMOFWSSTLPSKIOOWHOKOKNRKAQEDL

Important features of the protein: Signal peptide: amino acids 1-27

Transmembrane domain: amino acids 226-245

N-glycosylation site: amino acids 105-109

N-myristoylation sites: amino acids 10-16, 49-55, 62-68, 86-92, 150-156, 155-161, 162-168, 217-223, 227-233, 228-234, 232-238, 262-268, 357-363, 461-467

Prokaryotic membrane lipoprotein lipid attachment site: amino acids 12-23

Carboxylesterases type-B serine active site: amino acids 216-232

FIGURE 267

 $\tt TGTCGCCTGGCCCATGCAGACCCCGCGAGCGTCCCCTCCCGGCCCGGCCCTCCTG$ CTTCTGCTGCTGCTACTGGGGGGCGCCCACGGCCTCTTTCCTGAGGAGCCGCCGCCGCTT AGCGTGGCCCCCAGGGACTACCTGAACCACTATCCCGTGTTTGTGGGCAGCGGGCCCGGA CGCCTGACCCCCGCAGAAGGTGCTGACGACCTCAACATCCAGCGAGTCCTGCGGGTCAAC AGGACGCTGTTCATTGGGGACAGGGACAACCTCTACCGCGTAGAGCTGGAGCCCCCCACG TCCACGGAGCTGCGGTACCAGAGGAAGCTGACCTGGAGATCTAACCCCAGCGACATAAAC GTGTGTCGGATGAAGGGCAAACAGGAGGGCGAGTGTCGAAACTTCGTAAAGGTGCTGCTC CTTCGGGACGAGTCCACGCTCTTTGTGTGCGGTTCCAACGCCTTCAACCCGGTGTGCGCC AACTACAGCATAGACACCCTGCAGCCCGTCGGAGACAACATCAGCGGTATGGCCCGCTGC CCGTACGACCCCAAGCACGCCAATGTTGCCCTCTTCTCTGACGGGATGCTCTTCACAGCT ACTGTTACCGACTTCCTAGCCATTGATGCTGTCATCTACCGCAGCCTCGGGGACAGGCCC ACCCTGCGCACCGTGAAACATGACTCCAAGTGGTTCAAAGAGCCTTACTTTGTCCATGCG GTGGAGTGGGGCAGCCATGTCTACTTCTTCTCCGGGAGATTGCGATGGAGTTTAACTAC CTGGAGAAGGTGGTGTCCCGCGTGGCCCGAGTGTGCAAGAACGACGTGGGAGGCTCC CCCCGCGTGCTGGAGAAGCAGTGGACGTCCTTCCTGAAGGCGCGGCTCAACTGCTCTGTA CCCGGAGACTCCCATTTCTACTTCAACGTGCTGCAGGCTGTCACGGGCGTGGTCAGCCTC GGGGGCCGGCCCGTGGTCCTGGCCGTTTTTTCCACGCCCAGCAACAGCATCCCTGGCTCG GCTGTCTGCGCCTTTGACCTGACACAGGTGGCAGCTGTGTTTGAAGGCCGCTTCCGAGAG CAGAAGTCCCCCGAGTCCATCTGGACGCCGGTGCCGGAGGATCAGGTGCCTCGACCCCGG CCCGGGTGCTGCGCAGCCCCCGGGATGCAGTACAATGCCTCCAGCGCCTTGCCGGATGAC ATCCTCAACTTTGTCAAGACCCACCCTCTGATGGACGAGGCGGTGCCCTCGCTGGGCCAT GCGCCTGGATCCTGCGGACCCTGATGAGGCACCAGCTGACTCGAGTGGCTGTGGACGTG GGAGCCGGCCCCTGGGGCAACCAGACCGTTGTCTTCCTGGGTTCTGAGGCGGGGACGGTC CTCAAGTTCCTCGTCCGGCCCAATGCCAGCACCTCAGGGACGTCTGGGCTCAGTGTCTTC CTGGAGGAGTTTGAGACCTACCGGCCGGACAGGTGTGGACGGCCCGGCGGTGGCGAGACA GGGCAGCGGCTGCTGAGCTTGGAGCTGGACGCAGCTTCGGGGGGCCTGCTGGCTTC CCCCGCTGCGTGGTCCGAGTGCCTGTGGCTCCCAGCAGTACTCGGGGTGTATGAAG AACTGTATCGGCAGTCAGGACCCCTACTGCGGGTGGGCCCCCGACGGCTCCTGCATCTTC TTAGGGGACTGCACAGGACTCCTGCGGGCCAGCCTCTCCGAGGACCGCGCGGGGCTGGTG TCGGTGAACCTGCTGGTAACGTCGTCGGTGGCGGCCTTCGTGGTGGGAGCCGTGGTGTCC GGCTTCAGCGTGGGCTGGTTCGTGGGCCTCCGTGAGCGGCGGGAGCTGGCCCGGCGCAAG GACAAGGAGGCCATCCTGGCGCACGGGGCGGGCGAGGCGGTGCTGAGCGTCAGCCGCCTG GTTCCCCCGGAGGCCCTGCTGGCGCCCCTGATGCAGAACGGCTGGGCCAAGGCCACGCTG CTGCAGGGCGGGCCCCACGACCTGGACTCGGGGCTGCTGCCCACGCCCGAGCAGACGCCG CTGCCGCAGAAGCGCCTGCCCACTCCGCACCCGCACCCCCACGCCCTGGGCCCCCGCGCC CCCGCCGGGCCCCCGAGCAGCCCCCCGCGCCTGGGGAGCCGACCCCCGACGGCCGCCTC TATGCTGCCCGGCCCGGCCGCCCTCCCACGCGACTTCCCGCTCACCCCCCACGCCAGC CCGGACCGCCGGCGGTGTTCCGCGCCCACGGGCCCCTTGGACCCAGCCTCAGCCGCC GATGGCCTCCCGCGGCCCTGGAGCCCGCCCCGACGGGCAGCCTGAGGAGGCCACTGGGC CCCCACGCCCTCCGGCCGCCACCCTGCGCCGCACCCACACGTTCAACAGCGGCGAGGCC ATGCCTTGGCAGTGCCAGCCACGGGAACCAGGAGCGAGAGACGGTGCCAGAACGCCGGGG

FIGURE 268

MQTPRAS PPRPALLLLLLLGGAHGLFPEEPPLSVAPRDYLNHYPVFVGSGPGRLTPAE
GADDLNIGRVLRVNRTLFIGDRDNLYRVELEPPTSTELRYQRKLTWRSNPSDINVCRMKG
KQEGECRNFVKVLLIRDESTLFYCGSNAFNPVCANYSIDTLQPVGDNISGMARCPYDPKH
ANVALFSDGMLFTATVTDFLAIDAVIYRSLGDRPTLRTVKHDSKWFKEPYFVHAVEWGSH
VYFFFREIAMEFNYLEKVVVSRVARVCKNDVGGSPRVLEKQWTSFLKARLNCSVPGDSHF
YFNVLQAVTGVVSLGGRPVVLAVFSTENSIPGSAVCAFDLTQVAAVFEGRFREQKSPES
IWTPVPEDQVPRPRPGCCAAPGMQYNASSALPDDILMFVKTHPLMBEAVPSLGHAPWILR
TLMMHQLTRVAVDVGAGFWGNQTVVFLGSEAGTVLKFLVRPNASTSGTSGLSVFLEEFFT
YRPDRCGRPGGGETGQRLLSLELDAASGGLLAAFPRCVVRVPVARCQQYSGCMKNCTGSQ
DPYCGWAPDGSCIFLSPGTRAAFEQDVSGASTSGLGDCTGLLRASLSEDRAGLVSVNLLV
TSSVAAFVVGAVVSGFSVGWFVGLRERRELARRKDKEAILAHGAGEAVLSVSRLEGRRAQ
GPGGRGGGGGGGAGVPEPALLAPLMQNGWAKATLLQGGPHDLDSGLLPTPEGTPLPQKRL
PTPHPHPHALGPRAWDHGHPLLPASASSSLLLLAPARAPEQPPAGEPTPDGRLYAARPG
RASHGDFPLTPHASPDRRRVVSAPTGPLDPASAADGLPRPWSPPTGSLRRPLGPHAPPA
ATLRRTHTTNSGEARPGGRHRGCHARPGTDLAHLLPYGGADRTAPPVP

Important features of the protein: Signal peptide: amino acids 1-25

Transmembrane domains: amino acids 318-339, 598-617

441-445, 462-466

N-glycosylation sites. amino acids 74-78, 155-159, 167-171, 291-295, 386-390.

Glycosaminoglycan attachment sites: amino acids 51-55, 573-577

cAMP- and cGMP-dependent protein kinase phosphorylation site:

N-myristoylation sites: amino acids 21-27, 50-56, 189-195, 333-339, 382-388, 448-454, 490-496, 491-497, 508-514, 509-515, 531-537, 558-564, 569-575, 574-580, 580-586, 610-616, 643-649, 663-669, 666-672, 667-673, 668-674, 669-675, 670-676, 868-874, 879-885

FIGURE 269

ATCTGAGTGAGCTAACTGACACA<u>ATG</u>ANACTGTCAGGCATGTTTCTGCTCCTCTCTGG CTCTTTTCTGCTTTTTAACAGGTGTCTTCAGTCAGGGAGGACAGGTTGACTGGTGAGGT TCCAGGACCCCAAGGTCTACTGCACTCGGGAATCTAACCCACACTGTGGCTCTGATGGCC AGACATATGGCAATAAATGTGCCTTCTGTAAGGCCATAGTGAAAAGTGAGAAAGATTA GCCTAAAGCATCCTGGAAAATGCTGAGTTAAAGCCCATGTTTCTTGGTGACTTGCCAGCT TTTGCAGCCTTCTTTCTCACTTCTGCTTATACTTTTGCTGGTGATTCCTTTAATTCAT AAAGACATACCTACTCTGCCTGGGTCTTGAGGAGTTCAATGTATGCTATTTCTCTTGAT

FIGURE 270

MKLSGMFLLLSLALFCFLTGVFSQGGQVDCGEFQDPKVYCTRESNPHCGSDGQTYGNKCA FCKATVKSGGKISLKHPGKC

Important features of the protein: Signal peptide: amino acids 1-23

N-myristoylation sites: amino acids 26-32, 52-58, 56-62, 69-75

Kazal serine protease inhibitors family signature: amino acids 40-63

FIGURE 271

ACCATCATGTCCCTCCCACCCTCCTTTGACTGCGGGCCGTTCAGGTGCAGAGTCTCAGTT GCCCGGGAGCACCTCCCCTCCCGAGGCAGTCTGCTCAGAGGGCCTCGGCCCAGAATTCCA GTTCTGGTTTCATGCCAGCCTGTAAAAGGCCATGGAACTTTGGGTGAATCACCGATGCCA TTTAAGAGGGTTTTCTGCCAGGATGGAAATGTTAGGTCGTTCTGTGTCTGCGCTGTTCAT TTCAGTAGCCACCAGCCACCTGTGGCCGTTGAGTGCTTGAAATGAGGAACTGAGAAAATT AATTTCTCATGTATTTTCTCATTTATTTATTAATTTTTAACTGATAGTTGTACATATTT GGGGGTACATGTGATATTTGGATACATGTATACAATATATAATGATCAAATCAGGGTAAC TGGGATATCCATCACATCAAACATTTATTTTTTATTCTTTTTAGACAGAGTCTCACTCTG TCAAGCGATTCTCATGCCTCCACCTCCCAAGTAGCTGGGACTACAGGCATGCACCACAAT GCCCAACTAATTTTTGTATTTTTAGTAGAGACGGGGTTTTGCCATGTTGCCCAGGCTGGC CTTGAACTCCTGGCCTCAAACAATCCACTTGCCTCGGCCTCCCAAAGTGTTATGATTACA GGCGTGAGCCACCGTGCCTGGCCTAAACATTTATCTTTTCTTTGTGTTGGGAACTTTGAA ATTATACAATGAATTATTGTTAACTGTCATCTCCCTGCTGTGCTATGGAACACTGGGACT CTCTCTATCCTTCCCAACCTCTGATCACCTCATTCTACTCTCTACCTCCATGAGATCCAC TTTTTTAGCTCCCACATGTGAGTAAGAAAATGCAATATTTGTCTTTCTGTGCCTGGCTTA TTTCACTTAACATAATGACTTCCTGTTCCATCCATGTTGCTGCAAATGACAGGATTTCGT TCTTAATTTCAATTAAAATAACCACACATGGCAAAAA

FIGURE 272

 ${\tt MGLLLLVLFLSLLPVAYTIMSLPPSFDCGPFRCRVSVAREHLPSRGSLLRGPRPRIPVLV\\ {\tt SCQPVKGHGTLGESPMPFKRVFCQDGNVRSFCVCAVHFSSHQPPVAVECLK}$

Important features of the protein: Signal peptide: amino acids 1-18

N-myristoylation site: amino acids 86-92

Zinc carboxypeptidases, zinc-binding region 2 signature: amino acids 68-79

FIGURE 273

TTCTGAAGTAACGGAAGCTACCTTGTATAAAGACCTCAACACTGCTGACCATGATCAGCG CAGCCTGGAGCATCTTCCTCATCGGGACTAAAATTGGGCTGTTCCTTCAAGTAGCACCTC TATCAGTTATGGCTAAATCCTGTCCATCTGTGTGTCGCTGCGATGCGGGTTTCATTTACT GTAATGATCGCTTTCTGACATCCATTCCAACAGGAATACCAGAGGATGCTACAACTCTCT ACCTTCAGAACAACCAAATAAATAATGCTGGGATTCCTTCAGATTTGAAAAACTTGCTGA AAGTAGAAAGAATATACCTATACCACAACAGTTTAGATGAATTTCCTACCAACCTCCCAA AGTATGTAAAAGAGTTACATTTGCAAGAAAATAACATAAGGACTATCACTTATGATTCAC TTTCAAAAATTCCCTATCTGGAAGAATTACATTTAGATGACAACTCTGTCTCTGCAGTTA GCATAGAAGAGGGAGCATTCCGAGACAGCAACTATCTCCGACTGCTTTTCCTGTCCCGTA ATCACCTTAGCACAATTCCCTGGGGTTTGCCCAGGACTATAGAAGAACTACGCTTGGATG ATAATCGCATATCCACTATTTCATCACCATCTCTTCAAGGTCTCACTAGTCTAAAACGCC TGGTTCTAGATGGAAACCTGTTGAACAATCATGGTTTAGGTGACAAAGTTTTCTTCAACC TAGTTAATTTGACAGAGCTGTCCCTGGTGCGGAATTCCCTGACTGCTGCACCAGTAAACC TTCCAGGCACAAACCTGAGGAAGCTTTATCTTCAAGATAACCACATCAATCGGGTGCCCC CAAATGCTTTTTCTTATCTAAGGCAGCTCTATCGACTGGATATGTCCAATAATAACCTAA GTAATTTACCTCAGGGTATCTTTGATGATTTGGACAATATAACACAACTGATTCTTCGCA ACAATCCCTGGTATTGCGGGTGCAAGATGAAATGGGTACGTGACTGGTTACAATCACTAC CTGTGAAGGTCAACGTGCGTGGGCTCATGTGCCAAGCCCCAGAAAAGGTTCGTGGGATGG CTATTAAGGATCTCAATGCAGAACTGTTTGATTGTAAGGACAGTGGGATTGTAAGCACCA TTCAGATAACCACTGCAATACCCAACACAGTGTATCCTGCCCAAGGACAGTGGCCAGCTC CAGTGACCAAACAGCCAGATATTAAGAACCCCAAGCTCACTAAGGATCAACAAACCACAG GGAGTCCCTCAAGAAAAACAATTACAATTACTGTGAAGTCTGTCACCTCTGATACCATTC ATATCTCTTGGAAACTTGCTCTACCTATGACTGCTTTGAGACTCAGCTGGCTTAAACTGG GCCATAGCCCGGCATTTGGATCTATAACAGAAACAATTGTAACAGGGGAACGCAGTGAGT ACTTGGTCACAGCCCTGGAGCCTGATTCACCCTATAAAGTATGCATGGTTCCCATGGAAA CCAGCAACCTCTACCTATTTGATGAAACTCCTGTTTGTATTGAGACTGAAACTGCACCCC ACCCCAATTTACCTTTGGCTGCCATCATTGGTGGGGCTGTGGCCCTGGTTACCATTGCCC TTCTTGCTTTAGTGTTGGTATGTTCATAGGAATGGATCGCTCTTCTCAAGGAACTGTG CATATAGCAAAGGGAGGAGAAGAAAGGATGACTATGCAGAAGCTGGCACTAAGAAGGACA ACTCTATCCTGGAAATCAGGGAAACTTCTTTTCAGATGTTACCAATAAGCAATGAACCCA AAAACAATCACAGTGAAAGCAGTAGTAACCGAAGCTACAGAGACAGTGGTATTCCAGACT AAACCTAAGGGAGGTGATGGT

FIGURE 274

MISAAWSIFLIGTKIGLFLQVAPLSVMAKSCPSVCRCDAGFIYCNDRFLTSIPTGIPEDA
TTLYLQNNQINNAGIPSDLKNLLKVERIYLYHNSLDEFPTHLPKYVKELHLQENNIRTIT
YDSLSKIPYLEELHLDDNSVSAVSIEEGAFRDSNYLRLLFLSRNHLSTIPWGLPRTIEEL
RLDDNRISTISSPSLQGLTSLKRLVLDGNLLNHGLGGKVFFNLVNLTELSLVRNSLTAA
PVNLPGTNLRKLYLQDNHINRVPPNAFSYLRQLYRLDMSNNNLSNLPQGIFDDLDNITQL
ILRNNPWYCGCKMKWYRDWLQSLPVKVNVRGLMCQAPEKVRGMAIKDLNAELFDCKDGGI
VSTIQITTAIPNTVYPAQGQWPAPVTKQPDINNPKLTKDQQTTGSPSRKTTTITVKSVTS
DTIHISWKLALPMTALRSWLKLGHSPAFGSITETIVTGERSEYLVTALEPDSPYKVCMV
PMETSNLYLFDETPVCIETETAPLRMYNPTTTLNREQEKEPYKNPNLPLAAIIGGAVALV
TIALLALVCWYVHRNGSLFSRNCAYSKGRRKKDDYAEAGTKKDNSILBIRETSFQMLPIS
NEPISKEEFVIHTIFPPNGMNLYKNNHSESSSNRSYRDSGIPDSDHSHS

Important features of the protein: Signal peptide: amino acids 1-28

Transmembrane domain: amino acids 531-552

N-glycosylation sites: amino acids 226-229, 282-285, 296-299, 555-558, 626-629, 633-636

Tyrosine kinase phosphorylation site: amino acids 515-522

N-myristoylation sites: amino acids 12-17, 172-177, 208-213, 359-364, 534-539, 556-561. 640-645

Amidation site: amino acids 567-570

Leucine zipper pattern: amino acids 159-180

Phospholipase A2 aspartic acid active site: amino acids 34-44

FIGURE 275

AGGGCCGCGGGTGGAGAGAGCGACGCCCGAGGGGATGGCGCAGCGTCCCGGAGCGCCT TCTTCCAGCTGCAGCTGCAGGAGTTCATCAACGAGCGCGGCGTACTGGCCAGTGGGCGGC CTTGCGAGCCCGGCTGCCGGACTTTCTTCCGCGTCTGCCTTAAGCACTTCCAGGCGGTCG TCTCGCCCGGACCCTGCACCTTCGGGACCGTCTCCACGCCGGTATTGGGCACCAACTCCT TCGCTGTCCGGGACGACAGTAGCGGCGGGGGGGCGCAACCCTCTCCAACTGCCCTTCAATT TCACCTGGCCGGGTACCTTCTCGCTCATCATCGAAGCTTGGCACGCCCAGGAGACGACC TGCGGCCAGAGGCCTTGCCACCAGATGCACTCATCAGCAAGATCGCCATCCAGGGCTCCC TAGCTGTGGGTCAGAACTGGTTATTGGATGAGCAAACCAGCACCCTCACAAGGCTGCGCT ACTCTTACCGGGTCATCTGCAGTGACAACTACTATGGAGACAACTGCTCCCGCCTGTGCA TGCCCGGTTGGACTGGGGAATATTGCCAACAGCCTATCTGTCTTTCGGGCTGTCATGAAC AGAATGGCTACTGCAGCAAGCCAGCAGAGTGCCTCTGCCGCCCAGGCTGGCAGGGCCGGC TGTGTAACGAATGCATCCCCCACAATGGCTGTCGCCACGGCACCTGCAGCACTCCCTGGC AATGTACTTGTGATGAGGGCTGGGGAGGCCTGTTTTGTGACCAAGATCTCAACTACTGCA CCCACCACTCCCCATGCAAGAATGGGGCAACGTGCTCCAACAGTGGGCAGCGAAGCTACA CCTGCACCTGTCGCCCAGGCTACACTGGTGTGGACTGTGAGCTGGAGCTCAGCGAGTGTG ACAGCAACCCCTGTCGCAATGGAGGCAGCTGTAAGGACCAGGAGGATGGCTACCACTGCC TGTGTCCTCCGGGCTACTATGGCCTGCACTGTGAACACAGCACCTTGAGCTGCGCCGACT CCCCTGCTTCAATGGGGGCTCCTGCCGGGAGCGCAACCAGGGGGCCAACTATGCTTGTG AATGTCCCCCCAACTTCACCGGCTCCAACTGCGAGAAGAAGTGGACAGGTGCACCAGCA ACCCCTGTGCCAACGGGGGACAGTGCCTGAACCGAGGTCCAAGCCGCATGTGCCGCTGCC GTCCTGGATTCACGGGCACCTACTGTGAACTCCACGTCAGCGACTGTGCCCGTAACCCTT GCGCCCACGGTGGCACTTGCCATGACCTGGAGAATGGGCTCATGTGCACCTGCCCTGCCG GCTTCTCTGGCCGACGCTGTGAGGTGCGGACATCCATCGATGCCTGTGCCTCGAGTCCCT GCTTCAACAGGGCCACCTGCTACACCGACCTCTCCACAGACACCTTTGTGTGCAACTGCC CTTATGGCTTTGTGGGCAGCCGCTGCGAGTTCCCCGTGGGCTTGCCGCCCAGCTTCCCCT GGGTGGCCGTCTCGCTGGGTGTGGGGCTGCCAGTGCTGCTGGTACTGCTGGCATGGTGG CAGTGGCTGTGCGGCAGCTGCGGCTTCGACGGCCGGACGACGCAGCAGCAGCAAGCCATGA ACAACTTGTCGGACTTCCAGAAGGACAACCTGATTCCTGCCGCCCAGCTTAAAAACACAA ACCAGAGAGGAGCTGGAGTGGACTGTGGCCTGGACAAGTCCAACTGTGGCAAACAGC AAAACCACACTTGGACTATAATCTGGCCCCAGGGCCCCTGGGGCGGGGGACCATGCCAG GAAAGTTTCCCCACAGTGACAAGAGCTTAGGAGAGAGGCGCCACTGCGGTTACACAGTG AAAAGCCAGAGTGTCGGATATCAGCGATATGCTCCCCCAGGGACTCCATGTACCAGTCTG TGTGTTTGATATCAGAGGAGGAATGAATGTGTCATTGCCACGGAGGTA<u>TAA</u>GGCAGGA

FIGURE 276

MAAASRSASGWALLLLVALWQQRAAGSGVFQLQLQEFINERGVLASGRPCEPGCRTFFRV
CLKHFQAVVSPGPCTFGTVSTPVLGTINSFAVRDDSSGGRNPLQLPFNFTWPGTFSLIIE
AWHAPGDDLRPEALPPDALISKIAIQGSLAVGQNWLLDEQTSTLTRLRYSYRVICSDNYY
GDNCSRLCKKRNDHFGHYVCQPDGNLSCLPGWTGEYCQQPICLSGCHEQNGYCSKPAECL
CREGWQGRLCNECTPHNGCRHGTCSTPWQCTCDEGWGGLFCDQDLNYCTHHSPCKNGATC
SNSGQRSYTCTCRPGYTGVDCELELSECDSNPCRNGGSCKDQEDGYHCLCPPGYYGLHCE
HSTLSCADSPCFNGGSCRERNQGANYACECPPNFTGSNCEKKVDRCTSNPCANGGQCLNR
GPSRNCRCRPGFTGTYCELHVSDCARNPCAHGGTCHDLENGLMCTCPAGFSGRRCEVRTS
IDACASSPCFNRATCYTDLSTDTFVCNCPYGFVGSRCEFPVGLPPSFFWVAVSLGVGLAV
LLVLLGMVAVAVRQLRLRRPDDGSREAMNNLSDFQKDNLIPAAQLKNTNQKKELEVDCGL
DKSNCGKQQNHTLDYNLAPGPLGRGTMPGKFPHSDKSLGEKAPLRLHSEKPECRISAICS
PRDSMYQSVCLISEERNECVIATEV

Important features of the protein: Signal peptide: amino acids 1-26

Transmembrane domain: amino acids 530-552

N-glycosylation sites: amino acids 108-112, 183-187, 205-209, 393-397, 570-574, 610-614

Glycosaminoglycan attachment site: amino acids 96-100

Tyrosine kinase phosphorylation site: amino acids 340-347

N-myristoylation sites: amino acids 42-48, 204-210, 258-264, 277-283, 297-303, 383-389, 415-421, 461-467, 522-528, 535-541, 563-569, 599-605, 625-631

Amidation site: amino acids 471-475

Aspartic acid and asparagine hydroxylation site: amino acids 339-351

EGF-like domain cysteine pattern signature: amino acids 173-185, 206-218, 239-251, 270-282, 310-322, 348-360, 388-400, 426-438, 464-476, 506-518

Calcium-binding EGF-like: amino acids 224-245, 255-276, 295-316, 333-354, 373-394, 411-432, 449-470

FIGURE 277

TTTCGGTTTCTTCCAAGATTCCTGGCCTTCCCTCGACGGAGCCGGGCCCAGTGCGGGGGC GCAGGGGGGGGGGGGCTCCACCTCCTCGGCTTTCCCTGCGTCCAGAGGCTGGCATGGCGCG GGCCGAGTACTGAGCGCACGGTCGGGGCACAGCAGGGCCGGGGGGTGCAGCTGGCTCGCG CCTCCTCTCCGGCCGCCGTCTCCTCCGGTCCCTGGCGAAAGCCATTGAGACACCAGCTGG ACGTCACGCGCCGGAGCATGTCTGGGAGTCAGAGCGAGGTGGCTCCATCCCCGCAGAGTC $\tt CGCGGAGCCCCGAGATGGGACGGGACTTGCGGCCCGGGTCCCGCGTGCTCCTGCTCCTGC$ TTCTGCTCCTGCTGGTGTACCTGACTCAGCCAGGCAATGGCAACGAGGGCAGCGTCACTG GAAGTTGTTATTGTGGTAAAAGAATTTCTTCCGACTCCCCGCCATCGGTTCAGTTCATGA ATCGTCTCCGGAAACACCTGAGAGCTTACCATCGGTGTCTATACTACACGAGGTTCCAGC TCCTTTCCTGGAGCGTGTGTGGGGGCAACAAGGACCCATGGGTTCAGGAATTGATGAGCT GTCTTGATCTCAAAGAATGTGGACATGCTTACTCGGGGATTGTGGCCCACCAGAAGCATT TACTTCCTACCAGCCCCCCAATTTCTCAGGCCTCAGAGGGGGCATCTTCAGATATCCACA CCCCTGCCCAGATGCTCCTGTCCACCTTGCAGTCCACTCAGCGCCCCACCCTCCCAGTAG GATCACTGTCCTCGGACAAAGAGCTCACTCGTCCCAATGAAACCACCATTCACACTGCGG GCCACAGTCTGGCAGCTGGGCCTGAGGCTGGGGAGAACCAGAAGCAGCCGGAAAAAAATG CTGGTCCCACAGCCAGGACATCAGCCACAGTGCCAGTCCTGTGCCTCCTGGCCATCATCT TCATCCTCACCGCAGCCCTTTCCTATGTGCTGTGCAAGAGGAGGAGGGGGGCAGTCACCGC AGTCCTCTCCAGATCTGCCGGTTCATTATATACCTGTGGCACCTGACTCTAATACC<u>TGA</u>G CCAAGAATGGAAGCTTGTGAGGGTAAACTGTGGCTTATTCTTACAAAAAGTGTAATAAAG

FIGURE 278

MGRDLRPGSRVLLLLLLLLLVYLTQPGNGNEGSVTGSCYCGKRISSDSPPSVQFMNRLRK HLRAYHRCLYYTRFQLLSWSVCGGNKDPWVQELMSCLDLKECGHAYSGIVAHQKHLLPTS PPISQASEGASSDIHTPAQMLLSTLQSTQRPTLPVGSLSSDKELTRPNETTIHTAGHSLA AGPEAGENQKQPEKNAGPTARTSATVPVLCLLAIIFILTAALSYVLCKRRRGQSPQSSPD LPVHYIPVAPDSNT

Important features of the protein:

Signal peptide: 1-26

Transmembrane domain: 204-223

N-glycosylation site: 168-172

cAMP- and cGMP-dependent protein kinase phosphorylation site:

N-myristoylation site: 29-35, 32-38, 36-42, 156-162

Amidation site:

FIGURE 279

CGCGAGGCGCGGGAGCCTGGGACCAGGAGCGAGAGCCGCCTACCTGCAGCCGCCCCA CGGCACGGCAGCCACCATGCCGCTCCTGCTGTGCTTCGTGCTCCTGTGCGGAGTAGTGGA TTTCGCCAGAAGTTTGAGTATCACTACTCCTGAAGAGATGATTGAAAAAGCCAAAGGGGA AACTGCCTATCTGCCATGCAAATTTACGCTTAGTCCCGAAGACCAGGGACCGCTGGACAT CGAGTGGCTGATATCACCAGCTGATAATCAGAAGGTGGATCAAGTGATTATTTTATATTC TGGAGACAAAATTTATGATGACTACTATCCAGATCTGAAAGGCCGAGTACATTTTACGAG TAATGATCTCAAATCTGGTGATGCATCAATAAATGTAACGAATTTACAACTGTCAGATAT TGGCACATATCAGTGCAAAGTGAAAAAAGCTCCTGGTGTTGCAAATAAGAAGATTCATCT GGTAGTTCTTGTTAAGCCTTCAGGTGCGAGATGTTACGTTGATGGATCTGAAGAAATTGG AAGTGACTTTAAGATAAAATGTGAACCAAAAGAAGGTTCACTTCCATTACAGTATGAGTG GCAAAAATTGTCTGACTCACAGAAAATGCCCACTTCATGGTTAGCAGAAATGACTTCATC TGTTATATCTGTAAAAAATGCCTCTTCTGAGTACTCTGGGACATACAGCTGTACAGTCAG AAACAGAGTGGGCTCTGATCAGTGCCTGTTGCGTCTAAACGTTGTCCCTCCTTCAAATAA CGATATCAGGGAAGATGTGCCACCTCCAAAGAGCCGTACGTCCACTGCCAGAAGCTACAT CGGCAGTAATCATTCATCCCTGGGGTCCATGTCTCCTTCCAACATGGAAGGATATTCCAA GACTCAGTATAACCAAGTACCAAGTGAAGACTTTGAACGCACTCCTCAGAGTCCGACTCT CCCACCTGCTAAGTTCAAGTACCCTTACAAGACTGATGGAATTACAGTTGTATAAATATG GACTACTGAAGAATCTGAAGTATTGTATTATTTGACTTTATTTTAGGCCTCTAGTAAAGA CTTAAATGTTTTTTAAAAAAAGCACAAGGCACAGAGATTAGAGCAGCTGTAAGAACACAT CAAATTCTTTGTTAAAAAACCCTATGTATAGTGACACTGATAGTTAAAAGATGTTTTATT ATATTTTCAATAACTACCACTAACAAATTTTTAACTTTTCATATGCATATTCTGATATGT GGTCTTTTAGGAAAAGTATGGTTAATAGTTGATTTTTCAAAGGAAATTTTAAAATTCTTA CGTTCTGTTTAATGTTTTTGCTATTTAGTTAAATACATTGAAGGGAAATACCCGTTCTTT TCCCCTTTTATGCACACAACAGAAACACGCGTTGTCATGCCTCAAACTATTTTTATTTG CAACTACATGATTTCACACAATTCTCTTAAACAACGACATAAAATAGATTTCCTTGTATA TAAATAACTTACATACGCTCCATAAAGTAAATTCTCAAAGGTGCTAGAACAAATCGTCCA CTTCTACAGTGTTCTCGTATCCAACAGAGTTGATGCACAATATATAAATACTCAAGTCCA ATATTAAAAACTTAGGCACTTGACTAACTTTAATAAAATTTCTCAAACTATATCAATATC TAAAGTGCATATATTTTTTAAGAAAGATTATTCTCAATAACTTCTATAAAAATAAGTTTG ATGGTTTGGCCCATCTAACTTCACTACTATTAGTAAGAACTTTTAACTTTTAATGTGTAG TAAGGTTTATTCTACCTTTTTCTCAACATGACACCAACACAATCAAAAACGAAGTTAGTG AGGTGCTAACATGTGAGGATTAATCCAGTGATTCCGGTCACAATGCATTCCAGGAGGAGG AAATGGAACAGGAGGAGGATAGTGATTCTGATGGCCATTCCCTCGATACATTCCTGGCTT TTTTCTGGGCAAAGGGTGCCACATTGGAAGAGGTGGAAATATAAGTTCTGAAATCTGTAG GGAAGAGAACACATTAAGTTAATTCAAAGGAAAAAATCATCATCTATGTTCCAGATTTCT CATTAAAGACAAAGTTACCCACAACACTGAGATCACATCTAAGTGACACTCCTATTGTCA GGTCTAAATACATTAAAAACCTCATGTGTAATAGGCGTATAATGTATAACAGGTGACCAA TGTTTTCTGAATGCATAAAGAAATGAATAAACTCAAACACAGTACTTCCTAAACAACTTC AACCAAAAAAGACCAAAACATGGAACGAATGGAAGCTTGTAAGGACATGCTTGTTTAGT CCAGTGGTTTCCACAGCTGGCTAAGCCAGGAGTCACTTGGAGGCTTTTAAATACAAAACA TTGGAGCTGGAGGCCATTATCCTTAGCAAACTAATGCAGAAACAGAAAATCAACTACCGC ATGTTCTCACTTATAAGTGGGAGGTAATGATAAGAACTTATGAACACAAAGAAGGAAACA ATAGACATTGGAGTCTATTTGAGAGGGGAGGGTGGGAGAAGGAAAAGGAGCAGAAAAGAT AACTATTGAGTACTGCCTTCACACCTGGGTGATGAAATAATATGTACAACAAATCCCTGT GACACATGTTTACCTATGGAACAAACCTTCATGTGTATCCCTAAACCTAAAATAAAAGTT

FIGURE 280

MALLLCFVLLCGVVDFARSLSITTPEEMIEKAKGETAYLPCKFTLSPEDQGPLDIEWLIS
PADNOKVDOVITLYSGDKIYDDYYPDLKGRVHFTSNDLKSGDASINVTNLQLSDIGTYQG
KVKKAPQANKKHLVULVKPSGARCYVDGSEEIGSDFKIKCEPKEGSLPLQYEMQKLSD
SQKMPTSWLAEMTSSVISVKNASSEYSGTYSCTVRNRVGSDQCLLRLNVVPPSNKAGLIA
GAIIGTLLALALIGLIIFCCRKKRREEKYEKEVHHDIREDVPPPKSRTSTARSYIGSNHS
SLGSMSPSNMEGYSKTOYNQVPSEDFERTPQSPTLPPAKFKYPYKTDGITVV

Signal sequence.

Transmembrane domain: amino acids 236-257

N-glycosylation sites: amino acids 106-110, 201-205, 298-302

Tyrosine kinase phosphorylation sites: amino acids 31-39, 78-85, 262-270

N-myristoylation sites: amino acids 116-122, 208-214, 219-225, 237-243, 241-247, 245-251, 296-302

Myelin P0 protein: amino acids 96-125

FIGURE 281

TGCATCAGTGCCCAGGCAAGCCCAGGAGTTGACATTTCTCTGCCCAGCCATGGGCCTCAC CCTGCTCTTGCTGCTGCTCCTGGGACTAGAAGGTCAGGGCATAGTTGGCAGCCTCCCTGA GGTGCTGCAGGCACCCGTGGGAAGCTCCATTCTGGTGCAGTGCCACTACAGGCTCCAGGA TGTCAAAGCTCAGAAGGTGTGGTGCCGGTTCTTGCCGGAGGGGTGCCAGCCCCTGGTGTC CTCAGCTGTGGATCGCAGAGCTCCAGCGGGCAGGCGTACGTTTCTCACAGACCTGGGTGG GGGCCTGCTGCAGGTGGAAATGGTTACCCTGCAGGAAGAGGATGCTGGCGAGTATGGCTG CATGGTGGATGGGGCCAGGGGCCCCAGATTTTGCACAGAGTCTCTCTGAACATACTGCC CCCAGAGGAAGAAGAAGACCCATAAGATTGGCAGTCTGGCTGAGAACGCATTCTCAGA CCCTGCAGGCAGTGCCAACCCTTTGGAACCCAGCCAGGATGAGAAGAGCATCCCCTTGAT $\tt CTGGGGTGCTGCTCCTGGTAGGTCTGCTGGTGGCAGCGGTGGTGCTGTTTGCTGTGAT$ GGCCAAGAGGAAACAAGAATCCCTCCTCAGTGGTCCACCACGTCAG<u>TGA</u>CTCTGGACCGG ATACCACCTACACCAGCCTACCTCTTGATTCCCCATCAGGAAAACCTTCACTCCCAGCTC CATCCTCATTGCCCCCTCTACCTCCTAAGGTCCTGGTCTGCTCCAAGCCTGTGACATATG CCACAGTAATCTTCCCGGGAGGGAACAAGGGTGGAGGGACCTCGTGTGGGCCAGCCCAGA ATCCACCTAACAATCAGACTCCATCCAGCTAAGCTGCTCATCACACTTTAAACTCATGAG GACCATCCCTAGGGGTTCTGTGCATCCATCCAGCCAGCTCATGCCCTAGGATCCTTAGGA TATCTGAGCAACCAGGGACTTTAAGATCTAATCCAATGTCCTAACTTTACTAGGGAAAGT GACGCTCAGACATGACTGAGATGTCTTGGGGAAGACCTCCCTGCACCCAACTCCCCACT *ААААААААААААААААААААААААААААААА*

FIGURE 282

MGLTLLLLLLLGLEGQGIVGSLPEVLQAPVGSSILVQCHYRLQDVKAQKVWCRFLPEGCQ PLVSSAVDRRAPAGRRTFLTDLGGGLLQVEMVTLQEEDAGEYGCMVDGARGPQILHRVSL NILPPEEEETHKIGSLAENAFSDPAGSANPLEPSQDEKSIPLIWGAVLLVGLLVAAVVL FAVMAKRKOESLLSGPPRO

Important features of the protein:
Signal peptide:
amino acids 1-15

Transmembrane domain: amino acids 161-181

N-myristoylation sites: amino acids 17-23, 172-178

Amidation site: amino acids 73-79

FIGURE 283

FIGURE 284

MGLFMIIAILLFQKPTVTEQLKKCWNNYVQGHCRKICRVNEVPEALCENGRYCCLNIKEL EACKKITKPPRPKPATLALTLODYVTIIENFPSLKTQST

Important features of the protein:

Signal peptide:

Transmembrane domain:

cAMP- and cGMP-dependent protein kinase phosphorylation site:

FIGURE 285

GATGGCGCAGCCACAGCTTCTGTGAGATTCGATTTCTCCCCAGTTCCCCTGTGGGTCTGA GGGGACCAGAAGGGTGAGCTACGTTGGCTTTCTGGAAGGGGAGGCTATATGCGTCAATTC CCTGCTGTTCCAGGCCTTACCTGCTGGGCACTAACGGCGGAGCCAGGATGGGGACAGAAT AAAGGAGCCACGACCTGTGCCACCAACTCGCACTCAGACTCTGAACTCAGACCTGAAATC TTCTCTTCACGGGAGGCTTGGCAGTTTTTCTTACTCCTGTGGTCTCCAGATTTCAGGCCT AAGATGAAAGCCTCTAGTCTTGCCTTCAGCCTTCTCTCTGCTGCGTTTTATCTCCTATGG ACTCCTTCCACTGGACTGAAGACACTCAATTTGGGAAGCTGTGTGATCGCCACAAACCTT CAGGAAATACGAAATGGATTTTCTGAGATACGGGGCAGTGTGCAAGCCAAAGATGGAAAC ATTGACATCAGAATCTTAAGGAGGACTGAGTCTTTGCAAGACACAAAGCCTGCGAATCGA TGCTGCCTCCTGCGCCATTTGCTAAGACTCTATCTGGACAGGGTATTTAAAAACTACCAG ACCCCTGACCATTATACTCTCCGGAAGATCAGCAGCCTCGCCAATTCCTTTCTTACCATC AAGAAGGACCTCCGGCTCTCATGCCCACATGACATGCCATTGTGGGGAGGAAGCAATG AAGAAATACAGCCAGATTCTGAGTCACTTTGAAAAGCTGGAACCTCAGGCAGCAGTTGTG AGTGATGCTGCTAAGAATATTCGAGGTCAAGAGCTCCAGTCTTCAATACCTGCAGAG GAGGCATGACCCCAAACCACCATCTCTTTACTGTACTAGTCTTGTGCTGGTCACAGTGTA TCTTATTTATGCATTACTTGCTTCCTTGCATGATTGTCTTTATGCATCCCCAATCTTAAT CTTTAAAAAAATTCACAGATTATATTTATAACCTGACTAGAGCAGGTGATGTATTTTAT ACAGTAAAAAAAAAAACCTTGTAAATTCTAGAAGAGTGGCTAGGGGGGTTATTCATTTG TATTCAACTAAGGACATATTTACTCATGCTGATGCTCTGTGAGATATTTGAAATTGAACC AATGACTACTTAGGATGGGTTGTGGAATAAGTTTTGATGTGGAATTGCACATCTACCTTA AAAAAAAAA

FIGURE 286

MKASSLAFSLLSAAFYLLWTPSTGLKTLNLGSCVIATNLQEIRNGFSEIRGSVQAKDGNI DIRILRRTESLQDTKPANRCCLLRHLLRLYLDRVFKNYQTPDHYTLRKISSLANSFLTIK KDLRLSHAHMTCHCGEEAMKKYSQILSHFEKLEPQAAVVKALGELDILLQWMEETE

FIGURE 287

 ${\tt AATGCCCC} \underline{{\tt ATG}} {\tt CGCACCCCACAGCTCGCGCTCCTGCAAGTGTTCTTTCTGGTGTTCCCCG}$ $\overline{\texttt{ATGGCGTCCGGCCTCAGCCCTCTTCCTCCCCATCAGGGGCAGTGCCCACGTCTTTGGAGC}$ TGCAGCGAGGGACGGATGGCGGAACCCTCCAGTCCCCTTCAGAGGCGACTGCAACTCGCC CGGCCGTGCCTGGACTCCCTACAGTGGTCCCTACTCTCGTGACTCCCTCGGCCCCTGGGA ATAGGACTGTGGACCTCTTCCCAGTCTTACCGATCTGTGTCTGTGACTTGACTCCTGGAG CCTGCGATATAAATTGCTGCTGCGACAGGGACTGCTATCTTCTCCATCCGAGGACAGTTT TCTCCTTCTCCTTCCAGGCAGCGTAAGGTCTTCAAGCTGGGTTTGTGTAGACAACTCTG TTATCTTCAGGAGTAATTCCCCGTTTCCTTCAAGAGTTTTCATGGATTCTAATGGAATCA GGCAGTTTTGTGTCCATGTGAACAACTCAAACTTAAACTATTTCCAGAAGCTTCAAAAGG TCAATGCAACCAACTTCCAGGCCCTGGCTGCAGAGTTTGGAGGCGAATCATTCACTTCAA TCCCCAAGTGGTCTGTAATAAGCTTGCTGAGACAACCTGCAGGAGTTGGAGCTGGGGGAC TCTGTGCTGAAAGCAATCCTGCAGGTTTCCTAGAGAGTAAAAGTACAACTTGCACTCGTT TTTTCAAGAACCTGGCTAGTAGCTGTACCTTGGATTCAGCCCTCAATGCTGCCTCTTACT ATAACTTCACAGTCTTAAAGGTTCCAAGAAGCATGACTGATCCACAGAATATGGAGTTCC AGGTTCCTGTAATACTTACCTCACAGGCTAATGCTCCTCTGTTGGCTGGAAACACTTGTC AGAATGTAGTTTCTCAGGTCACCTATGAGATAGAGACCAATGGGACTTTTGGAATCCAGA AAGTTTCTGTCAGTTTGGGACAAACCAACCTGACTGTTGAGCCAGGCGCTTCCTTACAGC AACACTTCATCCTTCGCTTCAGGGCTTTTCAACAGAGCACAGCTGCTTCTCTCACCAGTC TAAGTTACTCAATGACCCTCTTACAGAGCCAGGGTAATGGAAGTTGCTCTGTTAAAAGAC ATGAAGTGCAGTTTGGAGTGAATGCAATATCTGGATGCAAGCTCAGGTTGAAGAAGGCAG ACTGCAGCCACTTGCAGCAGGAGATTTATCAGACTCTTCATGGAAGGCCCAGACCAGAGT ATGTTGCCATCTTTGGTAATGCTGACCCAGCCCAGAAAGGAGGGTGGACCAGGATCCTCA ACAGGCACTGCAGCATTTCAGCTATAAACTGTACTTCCTGCTGTCTCATACCAGTTTCCC TGGAGATCCAGGTATTGTGGGCATATGTAGGTCTCCTGTCCAACCCGCAAGCTCATGTAT CAGGAGTTCGATTCCTATACCAGTGCCAGTCTATACAGGATTCTCAGCAAGTTACAGAAG TATCTTTGACAACTCTTGTGAACTTTGTGGACATTACCCAGAAGCCACAGCCTCCAAGGG GCCAACCCAAAATGGACTGGAAATGGCCATTCGACTTCTTTCCCTTCAAAGTGGCATTCA GCAGAGGAGTATTCTCTCAAAAATGCTCAGTCTCTCCCATCCTTATCCTGTGCCTCTTAC TACTTGGAGTTCTCAACCTAGAGACTATGTGAAGAAAAGAAAATAATCAGATTTCAGTTT TCCCTATGAGAAACTCTGAGGCAGCCACTTATCTTGGCTAAATAGAACCTCACCTGCTCA TGACCAGAGAGCATTTAGGATAATAGATGACCTAACTGAAGGAATCCTTGTATATGAAAG

FIGURE 288

MRTPQLALLQVFFLVFPDGVRPQPSSSPSGAVPTSLELQRGTDGGTLQSPSEATATRPAV
PGLPTVVPTLVTPSAPENRTVDLFPVLPICVCDLTPGACDINCCCDRDCYLLHPRTVFSF
CLPGSVRSSSWVCVDNSVIFRSNSPFPSRVFMDSNGIRQFCVHVNNSNLNYFQKLQKVNA
TNFQALAAEFGGESFTSTFQTQSPPSFYRAGDPILTYFPKWSVISLLRQPAGVGAGGLC
ESNPAGFLESKSTTCTRFFKNLASSCTLDSALNAASYYNFTVLKVPRSMTDPONMEFQVP
VILTSQANAPLLAENTCQNVVSQVTYEIETNGTFGIQKVSVSLGQTNLTVEPGASLQQHF
ILRFRAFQQSTAASLTSPRSGNPGYIVGKPLLALTDDISYSMTLLQSQGNGSCSVKRHEV
OFGVNAISGCKLRLKKADCSHLQGEIYGTLHGRPRPEVAIFGNADPAQKGGWTRILNRH
CSISAINCTSCCLIPVSLEIQVLWAYVGLLSNPQAHVSGVRFLYQCQSIQDSQQVTEVSL
TTLVNFVDITQKPQPPRGQPKMDWKWPPDFFPFKVAFSRGVFSQKCSVSPILILCLLLLG
VINNEYM

Important features of the protein: Signal peptide: amino acids 1-22

Transmembrane domains: amino acids 484-505, 581-600

N-glycosylation sites: amino acids 78-82, 165-169, 179-185, 279-285, 331-337, 347-351, 410-414, 487-491

N-myristoylation sites: amino acids 30-36, 41-47, 124-130, 232-238, 236-242, 409-415

Prokaryotic membrane lipoprotein lipid attachment site: amino acids 420-431

FIGURE 289

CGCGGAGCCCTGCGCTGGGAGGTGCACGGTGTGCACGCTGGACTGGACCCCCATGCAACC CCGCGCCCTGCGCCTTAACCAGGACTGCTCCGCGCGCCCCTGAGCCTCGGGCTCCGGCCC AAAGATGGCTTTAAAAGTGCTACTAGAACAAGAGAAAACGTTTTTCACTCTTTTAGTATT ACTAGGCTATTTGTCATGTAAAGTGACTTGTGAATCAGGAGACTGTAGACAGCAAGAATT CAGGGATCGGTCTGGAAACTGTGTTCCCTGCAACCAGTGTGGGCCAGGCATGGAGTTGTC TAAGGAATGTGGCTTCGGCTATGGGGAGGATGCACAGTGTGTGACGTGCCGGCTGCACAG GTTCAAGGAGGACTGGGGCTTCCAGAAATGCAAGCCCTGTCTGGACTGCGCAGTGGTGAA CCGCTTTCAGAAGGCAAATTGTTCAGCCACCAGTGATGCCATCTGCGGGGACTGCTTGCC AGGATTTTATAGGAAGACGAAACTTGTCGGCTTTCAAGACATGGAGTGTGTGCCTTGTGG AGACCCTCCTCCTCCTTACGAACCGCACTGTGCCAGCAAGGTCAACCTCGTGAAGATCGC GTCCACGGCCTCCAGCCCACGGGACACGGCGCTGGCTGCCGTTATCTGCAGCGCTCTGGC CACCGTCCTGCTGGCCCTGCTCATCCTCTGTGTCATCTATTGTAAGAGACAGTTTATGGA GAAGAAACCCAGCTGGTCTCTGCGGTCGCAGGACATTCAGTACAACGGCTCTGAGCTGTC GTGTTTTGACAGACCTCAGCTCCACGAATATGCCCACAGAGCCTGCTGCCAGTGCCGCCG CTGCAGCCCCAACCCGGCGACTCTTGGTTGTGGGGTGCATTCTGCAGCCAGTCTTCAGGC AAGAAACGCAGGCCCAGCCGGGGAGATGGTGCCGACTTTCTTCGGATCCCTCACGCAGTC CATCTGTGGCGAGTTTTCAGATGCCTGGCCTCTGATGCAGAATCCCATGGGTGGTGACAA TGTTCCAGTCCAGTCTCATTCTGAAAACTTTACAGCAGCTACTGATTTATCTAGATATAA TCAGGAGAGTGGCGCTGTCATCCACCCAGCCACTCAGACGTCCCTCCAGGAAGCT<u>TAA</u>AG AACCTGCTTCTTCTGCAGTAGAAGCGTGTGCTGGAACCCAAAGAGTACTCCTTTGTTAG AAACTGACGGCATTTGAAGCCTTTCAGCCAGTTGCTTCTGAGCCAGACCAGCTGTAAGCT GAAACCTCAATGAATAACAAGAAAAGACTCCAGGCCGACTCATGATACTCTGCATCTTTC CTACATGAGAAGCTTCTCTGCCACAAAAGTGACTTCAAAGACTGATGGGTTGAGCTGGCA GCCTATGAGATTGTGGACATATAACAAGAAACAGAAATGCCCTCATGCTTATTTTCATGG CATACCGCCTATGAAATATCAGATAAATTACCTTAGCTTTTATGTAGAATGGGTTCAAAA GTGAGTGTTTCTATTTGAGAAGGACACTTTTTCATCATCTAAACTGATTCGCATAGGTGG TTAGAATGGCCCTCATATTGCCTGCCTAAATCTTGGGTTTATTAGATGAAGTTTACTGAA TCAGAGGAATCAGACAGAGGAGGATAGCTCTTTCCAGAATCCACACTTCTGACCTCAGCC TCGGTCTCATGAACACCCGCTGATCTCAGGAGAACACCTGGGCTAGGGAATGTGGTCGAG AAAGGGCAGCCCATTGCCCAGAATTAACACATATTGTAGAGACTTGTATGCAAAGGTTGG CATATTTATATGAAAATTAGTTGCTATAGAAACATTTGTTGCATCTGTCCCTCTGCCTGA GCTTAGAAGGTTATAGAAAAAGGGTATTTATAAACATAAATGACCTTTTACTTGCATTGT ATCTTATACTAAAGGCTTTAGAAATTACAACATATCAGGTTCCCCTACTACTGAAGTAGC CTTCCGTGAGAACACCACCACGTTAGGACTAGAAGAAAATGCACAATTTGTAGGGGTTT GGATGAAGCAGCTGTAACTGCCCTAGTGTAGTTTGACCAGGACATTGTCGTGCTCCTTCC AATTGTGTAAGATTAGTTAGCACATCATCTCCTACTTTAGCCATCCGGTGTTGGATTTAA GAGGACGGTGCTTCTTTCTATTAAAGTGCTCCATCCCCTACCATCTACACATTAGCATTG ${\tt TCTCTAGAGCTAAGACAGAAATTAACCCCGTTCAGTCACAAAGCAGGGAATGGTTCATTT}$ ACTCTTAATCTTTATGCCCTGGAGAAGACCTACTTGAACAGGGCATATTTTTTAGACTTC TGAACATCAGTATGTTCGAGGGTACTATGATATTTTGGTTTTGGAATTGCCCTGCCCAAGT CACTGTCTTTTAACTTTTAAACTGAATATTAAAATGTATCTGTCTTTCCT

FIGURE 290

MALKVLLEQEKTFFTLLVLLGYLSCKVTCESGDCRQQEFRDRSGNCVPCNQCGPGMELSK ECGFGYGEDAQCVTCRLHRFKEDWGFQKCKPCLDCAVVNRFQKANCSATSDAICGDCLPG FYRKTKLVGFQDMECVPCGDPPPYEPHCASKVNLVKIASTASSPRDTALAAVICSALAT VLLALLILCVIYCKRQFMEKKPSWSLRSQDIQYNGSELSCFDRPQLHEYAHRACCQCRRD SVQTCGPVRLLPSMCCEEACSPNPATLGCGVHSAASLQARNAGPAGEMVPTFFGSLTQSI CGEFSDAWPLMQNPMGGDNISFCDSYPELTGEDIHSLNPELESSTSLDSNSSQUYEQGAV PVQSHSENFTAATDLSRYNNTLVESASTQDALTMRSQLDQESGAVIHPATQTSLQEA

Important features of the protein: Signal peptide: Amino acids1-25

Transmembrane domain:

Amino acids 169-192

N-glycosylation sites: Amino acids105-109;214-218;319-323;350-354;368-372;379-383

cAMP- and cGMP-dependent protein kinase phosphorylation sites:
Amino acids 200-204:238-242

Tyrosine kinase phosphorylation site: Amino acids 207-214

N-myristoylation sites: Amino acids 55-61; 215-221; 270-276

Prokaryotic membrane lipoprotein lipid attachment site: Amino acids 259-270

TNFR/NGFR family cysteine-rich region proteins: Amino acids 89-96

FIGURE 291

CCTGGAGCCGGAAGCGCGGCTGCAGCAGGGCGAGGCTCCAGGTGGGGTCGGTTCCGCATC CAGCCTAGCGTGTCCACGATGCGGCTGGGCTCCGGGACTTTCGCTACCTGTTGCGTAGCG TCTGCCAGAGCGGAACACGGAGCGGAGCCCCCAGCGCCCGAACCCTCGGCTGGAGCCAGT TCTAACTGGACCACGCTGCCACCACCTCTCTTCAGTAAAGTTGTTATTGTTCTGATAGAT GCCTTGAGAGATGATTTTGTGTTTGGGTCAAAGGGTGTGAAATTTATGCCCTACACAACT TACCTTGTGGAAAAAGGAGCATCTCACAGTTTTGTGGCTGAAGCAAAGCCACCTACAGTT ACTATGCCTCGAATCAAGGCATTGATGACGGGGAGCCTTCCTGGCTTTGTCGACGTCATC AGGAACCTCAATTCTCCTGCACTGCTGGAAGACAGTGTGATAAGACAAGCAAAAGCAGCT GGAAAAAGAATAGTCTTTTATGGAGATGAAACCTGGGTTAAATTATTCCCAAAGCATTTT GTGGAATATGATGGAACAACCTCATTTTTCGTGTCAGATTACACAGAGGTGGATAATAAT GTCACGAGGCATTTGGATAAAGTATTAAAAAGAGGAGATTGGGACATATTAATCCTCCAC TACCTGGGGCTGGACCACATTGGCCACATTTCAGGGCCCAACAGCCCCCTGATTGGGCAG AAGCTGAGCGAGATGGACAGCGTGCTGATGAAGATCCACACCTCACTGCAGTCGAAGGAG AGAGAGACGCCTTTACCCAATTTGCTGGTTCTTTGTGGTGACCATGGCATGTCTGAAACA GGAAGTCACGGGGCCTCCTCCACCGAGGAGGTGAATACACCTCTGATTTTAATCAGTTCT GCGTTTGAAAGGAAACCCGGTGATATCCGACATCCAAAGCACGTCCAA<u>TAG</u>ACGGATGTG GCTGCGACACTGGCGATAGCACTTGGCTTACCGATTCCAAAAGACAGTGTAGGGAGCCTC CTATTCCCAGTTGTGGAAGGAAGACCAATGAGAGAGCAGTTGAGATTTTTACATTTGAAT ACAGTGCAGCTTAGTAAACTGTTGCAAGAGAATGTGCCGTCATATGAAAAAGATCCTGGG TTTGAGCAGTTTAAAATGTCAGAAAGATTGCATGGGAACTGGATCAGACTGTACTTGGAG GAAAAGCATTCAGAAGTCCTATTCAACCTGGGCTCCAAGGTTCTCAGGCAGTACCTGGAT GCTCTGAAGACGCTGAGCTTGTCCCTGAGTGCACAAGTGGCCCAGTTCTCACCCTGCTCC TGCTCAGCGTCCCACAGGCACTGCACAGAAAGGCTGAGCTGGAAGTCCCACTGTCATCTC $\tt CTGGGTTTTCTCTGCTCTTTTATTTGGTGATCCTGGTTCTTTCGGCCGTTCACGTCATTG$ GCCTTTCGTTTACCAGACTCTGGTTGAACACCTGGTGTGTGCCAAGTGCTGGCAGTGCCC TGGACAGGGGCCTCAGGGAAGGACGTGGAGCAGCCTTATCCCAGGCCTCTGGGTGTCCC GACACAGGTGTTCACATCTGTGCTGTCAGGTCAGATGCCTCAGTTCTTGGAAAGCTAGGT TCCTGCGACTGTTACCAAGGTGATTGTAAAGAGCTGGCGGTCACAGAGGAACAAGCCCCC CAGCTGAGGGGGTGTGTGAATCGGACAGCCTCCCAGCAGAGGTGTGGGAGCTGCAGCTGA GGGAAGAAGAGACAATCGGCCTGGACACTCAGGAGGGTCAAAAGGAGACTTGGTCGCACC CGGACGTTTTCTGTTGGAATTCTTAGTCCTTGGCCTCGGACACCTTCATTCGTTAGCTGG GGAGTGGTGGTGAGGCAGTGAAGAAGAGGCGGATGGTCACACTCAGATCCACAGAGCCCA GGATCAAGGGACCCACTGCAGTGGCAGCAGGACTGTTGGGCCCCCACCCCAACCCTGCAC AGCCCTCATCCCCTCTTGGCTTGAGCCGTCAGAGGCCCTGTGCTGAGTGTCTGACCGAGA CACTCACAGCTTTGTCATCAGGGCACAGGCTTCCTCGGAGCCAGGATGATCTGTGCCACG CTTGCACCTCGGGCCCATCTGGGCTCATGCTCTCTCTCTGCTATTGAATTAGTACCTAG

FIGURE 292

MRLGSGTFATCCVAIEVLGIAVFLRGFFPAPVRSSARAEHGAEPPAPEPSAGASSNWTTL
PPPLFSKVVIVLIDALRDDFVFGSKGVKFMPYTTTLVEKGASHSFVAEAKPPTVTMPRIK
ALMTGSLPGFVDVIRNLNSPALLEDSVIRQAKAAGKRIVFYGDETWVKLFFKHFVEYDGT
TSFFVSDYTEVDNNVTRHLDKVLKRGDWDILIILHYLGLDHIGHISGPNSPLIGQKLSEMD
SVLMKIHTSLQSKERETPLPNLLVLCGDHGMSETGSHGASSTEEVNTPLILISSAFERKP
GDIRHPKHVO

Important features of the protein: Signal peptide: amino acids 1-34

Transmembrane domain: amino acids 58-76

N-glycosylation sites: amino acids 56-60, 194-198

N-myristoylation sites: amino acids 6-12, 52-58, 100-106, 125-131, 233-239, 270-276, 275-281, 278-284

Amidation site: amino acids 154-158

Cell attachment sequence: amino acids 205-208

FIGURE 293

AGCCAGGCAGCACATCACAGCGGGAGGAGCTGTCCCAGGTGGCCCAGCTCAGCAATGGCA ATGGGGGTCCCCAGAGTCATTCTGCTCTGCCTCTTTGGGGCTGCCTCTGCCTGACAGGG TCCCAAGCCCTGCAGTGCTACAGCTTTGAGCACACCTACTTTGGCCCCCTTTGACCTCAGG GCCATGAAGCTGCCCAGCATCTCCTGTCCTCATGAGTGCTTTGAGGCTATCCTGTCTCTG GACACCGGGTATCGCGCGCCGGTGACCCTGGTGCGGAAGGGCTGCTGGACCGGGCCTCCT GCGGGCCAGACGCAATCGAACCCGGACGCGCTGCCGCCAGACTACTCGGTGGTGCGCGGC TGCACAACTGACAAATGCAACGCCCACCTCATGACTCATGACGCCCTCCCCAACCTGAGC CAAGCACCCGACCCGCCGACGCTCAGCGGCGCCGAGTGCTACGCCTGTATCGGGGTCCAC CAGGATGACTGCGCTATCGGCAGGTCCCGACGAGTCCAGTGTCACCAGGACCAGACCGCC TGCTTCCAGGGCAGTGGCAGATGACAGTTGGCAATTTCTCAGTCCCTGTGTACATCAGA ACCTGCCACCGGCCCTCCTGCACCACCGAGGGCACCACCAGCCCCTGGACAGCCATCGAC CTCCAGGGCTCCTGCTGTGAGGGGTACCTCTGCAACAGGAAATCCATGACCCAGCCCTTC ACCAGTGCTTCAGCCACCACCCCTCCCCGAGCACTACAGGTCCTGGCCCTGCTCCTCCCA $\tt GTCCTCCTGCTGGTGGGGCTCTCAGCA{\color{red}{TAG}}ACCGCCCCTCCAGGATGCTGGGGACAGGGC$ TCACACACCTCATTCTTGCTGCTTCAGCCCCTATCACATAGCTCACTGGAAAATGATGTT AAAGTAAGAATTGCAAAA

FIGURE 294

MAMGVPRVILLCLFGAALCLTGSQALQCYSFEHTYFGPFDLRAMKLPSISCPHECFEAIL SLDTGYRAPVTLVRKGCWTGPPAGQTQSNPDALPPDYSVVRGCTTDKCNAHLMTHDALPN LSQAPDPPTLSGAECYACIGVHQDDCAIGRSRRVQCHQDQTACFQGSGRMTVGNFSVPVY IRTCHRPSCTTEGTTSPWTAIDLQGSCCEGYLCNRKSMTQPFTSASATTPPRALQVLALL LPVLLLVGLSA

Important features of the protein: Signal peptide: amino acids 1-19

Transmembrane domain: amino acids 233-251

N-glycosylation sites: amino acids 120-124, 174-178

N-myristoylation sites: amino acids 15-21, 84-90

FIGURE 295

FIGURE 296

MVMLLLLLSALAGLFGAAEGQAFHLGKCPNPPVQENFDVNKYLGRWYEIEKIPTTFENG RCIQANYSLMENGKIKVLNQELRADGTVNQIEGEATPVNLTEPAKLEVKFSWFMPSAPY WILATDYENYALVYSCTCIIQLFHVDFAWILARNPNLPPETVDSLKNILTSNNIDVKKM TVTDOVNCPKLS

Signal sequence:

1-16

N-glycosylation site:

65-68

98-101

cAMP- and cGMP-dependent protein kinase phosphorylation

site: 175-178

N-myristoylation site:

13-18

16-21

Lipocalin proteins:

36-47

120-130

Lipocalin / cytosolic fatty-acid binding proteins:

41-185

FIGURE 297

GGGTGATTGAACTAAACCTTCGCCGCACCGAGTTTGCAGTACGGCCGTCACCCGCACCGC TGCCTGCTTGCGGTTGGAGAAATCAAGGCCCTACCGGGCCTCCGTAGTCACCTCTATA GTGGGCGTGGCCGAGGCCGGGTGACCCTGCCGGAGCCTCCGCTGCCAGCGACATGTTCA AGGTAATTCAGAGGTCCGTGGGGCCAGCCAGCCTGAGCTTGCTCACCTTCAAAGTCTATG CAGCACCAAAAAAGGACTCACCTCCCAAAAATTCCGTGAAGGTTGATGAGCTTTCACTCT ACTCAGTTCCTGAGGGTCAATCGAAGTATGTGGAGGAGGCAAGGAGCCAGCTTGAAGAAA GCATCTCACAGCTCCGACACTATTGCGAGCCATACACAACCTGGTGTCAGGAAACGTACT CCCAAACTAAGCCCAAGATGCAAAGTTTGGTTCAATGGGGGTTAGACAGCTATGACTATC TCCAAAATGCACCTCCTGGATTTTTTCCGAGACTTGGTGTTATTGGTTTTTGCTGGCCTTA TTGGACTCCTTTTGGCTAGAGGTTCAAAAATAAAGAAGCTAGTGTATCCGCCTGGTTTCA TGGGATTAGCTGCCTCCTCTATTATCCACAACAACCCATCGTGTTTTGCCCAGGTCAGTG CCCACAGATTA TATGACTGGGGTTTACGAGGATATATAGTCATAGAAGATTTGTGGAAGG AGAACTTTCAAAAGCCAGGAAATGTGAAGAATTCACCTGGAACTAAGTAGAAAACTCCAT GCTCTGCCATCTTAATCAGTTATAGGTAAACATTGGAAACTCCATAGAATAAATCAGTAT TTCTACAGAAAATGGCATAGAAGTCAGTATTGAATGTATTAAATTGGCTTTCTTCTTCA GGAAAAACTAGACCAGACCTCTGTTATCTTCTGTGAAATCATCCTACAAGCAAACTAACC TGGAATCCCTTCACCTAGAGATAATGTACAAGCCTTAGAACTCCTCATTCTCATGTTGCT AAAAAAA

FIGURE 298

MFKVIQRSVGPASLSLLTFKVYAAPKKDSPPKNSVKVDELSLYSVPEGQSKYVEEARSQL EESISQLRHYCEPYTTWCQETYSQTKPKMQSLVQWGLDSYDYLQNAPPGFFFRLGVIGFA GLIGLLLARGSKIKKLVYPPGFMGLAASLYYPQQAIVFAQVSGERLYDWGLRGYIVIEDL WKENFOKPGNVKNSPGTK

Important features: Signal peptide: Amino acids 1-23

Transmembrane domain: . Amino acids 111-130

 $\mathtt{cAMP-}$ and $\mathtt{cGMP-}dependent$ protein kinase phosphorylation site: Amino acids 26-30

Tyrosine kinase phosphorylation site: Amino acids 36-44

N-myristoylation sites: Amino acids 124-130;144-150;189-195

FIGURE 299

CCGCTGAGATGTACGAACTTCCGGTTCTCCGGGCAGCTGCCACTGCTGTAGCTTCTGCCA CCTGCCACGACCGGGCCTCTCCCTGGCGTTTGGTCACCTCTGCTTCATTCTCCACCGCGC CTATGGTCCCTCTTGGAGCCAGCGTGGCGGGCCTGGCGGCTCCCGGGTGGTGAGAGAGCG GTCCGGGAACGATGAAGGCCTCGCAGTGCTGCTGTCTCAGCCACCTCTTGGCTTCCG TCCTCCTCCTCCTCCTCCTCCTCAACTAACCGCCCCCTGGCAGTCCTGCTGCAGCAG $\tt CCGAGGCCGCGCCAGGTCTTGGGCCTCCTGACCCTAGACCACGGACATTACCGCCGCTGC$ CACCGGGCCCTACCCCTGCCCAGCAGCCGGGCCGTGGTCTGGCTGAAGCTGCGGGGCCGC GAGGGAAGGCCGGGGAAGGCTCGGTGGGTGGCGCCCTTGCTGTGAGCCCCAACCCTGGCG ACAAGCCCATGACCCAGCGGGCCCTGACCGTGTTGATGGTGATGAGCGGCGCGGTGCTGG TGTACTTCGTGGTCAGGACGGTCAGGATGAGAAGAAGAAACCGAAAGACTAGGAGATATG GAGTTTTGGACACTAACATAGAAAATATGGAATTGACACCTTTAGAACAGGATGATGAGG ATGATGACAACACGTTGTTTGATGCCAATCATCCTCGAAGA**TAA**GAATGTGCCTTTTGAT GAAAGAACTTTATCTTTCTACAATGAAGAGTGGAATTTCTATGTTTAAGGAATAAGAAGC CACTATATCAATGTTGGGGGGGTATTTAAGTTACATATATTTTAACAACCTTTAATTTGC TGTTGCAATAAATACCGTATCCTTTTATTATATCTTTATATGTATAGAAGTACTCTATTA ATGGGCTCAGAGATGTTGGGGATAAAGTATACTGTAATAATTTATCTGTTTGAAAATTAC TATAAAACGGTGTTTTCTGGTCGGTTTTTGTTTCCTGCTTACCATATGATTGTAAATTGT TTTATGTATTAATCAGTTAATGCTAATTATTTTTGCTGATGTCATATGTTAAAGAGCTAT AAATTCCAACAACCAACTGGTGTAAAAATAATTTAAAATTTCCTTTACTGAAAGGTAT TTCCCATTTTTGTGGGGAAAAGAAGCCAAATTTATTACTTTGTGTTGGGGTTTTTAAAAT ATTA AGA A TGTCTA AGTTATTGTTTGCAAAACAATAAATATGATTTTAAATTCTCTTAA AAAAAAA

FIGURE 300

MKASQCCCLSHLLASVLLLLLLPELSGPLAVLLQAAEAAPGLGPPDPRPRTLPPLPPGP TPAQQPGRGLAEAAGPRGSEGGNGSNPVAGLETDDHGGKAGEGSVGGGLAVSPNPGDKPM TQRALTVLMVVSGAVLVYFVVRTVRMRRRNRKTRRYGVLDTNIENMELTPLEQDDEDDDN TLFDANHPRR

Signal peptide: amino acids 1-28

Transmembrane domain: amino acids 124-140

N-glycosylation site: amino acids 83-87

N-myristoylation sites: amino acids 69-75, 78-84, 81-87, 97-103, 103-109, 106-112, 157-160

FIGURE 301

CTCGGCTGGATTTAAGGTTGCCGCTAGCCGCCTGGGAATTTAAGGGACCCACACTACCTT CCCGAAGTTGAAGGCAAGCGGTGATTGTTTGTAGACGGCGCTTTGTCATGGGACCTGTGC GGTTGGGAATATTGCTTTTCCTTTTTTTGGCCGTGCACGAGGCTTGGGCTGGGATGTTGA AGGAGGAGGACGATGACACAGAACGCTTGCCCAGCAAATGCGAAGTGTGTAAGCTGCTGA GCACAGAGCTACAGGCGGAACTGAGTCGCACCGGTCGATCTCGAGAGGTGCTGGAGCTGG GGCAGGTGCTGGATACAGGCAAGAGGAAGAGACACGTGCCTTACAGCGTTTCAGAGACAA GGCTGGAAGAGGCCTTAGAGAATTTATGTGAGCGGATCCTGGACTATAGTGTTCACGCTG AGCGCAAGGGCTCACTGAGATATGCCAAGGGTCAGAGTCAGACCATGGCAACACTGAAAG GCCTAGTGCAGAAGGGGGTGAAGGTGGATCTGGGGATCCCTCTGGAGCTTTGGGATGAGC CCAGCGTGGAGGTCACATACCTCAAGAAGCAGTGTGAGACCATGTTGGAGGAGTTTGAAG ACATTGTGGGAGACTGGTACTTCCACCATCAGGAGCAGCCCCTACAAAATTTTCTCTGTG AAGGTCATGTGCTCCCAGCTGCTGAAACTGCATGTCTACAGGAAACTTGGACTGGAAAGG AGGAGGAAGAGGAAGGGGAGACAAGATGACCAAGACAGGAAGCCACCCCAAACTTGACC TCTAAAGCCTGCACTCTCCCTGCTCCACAGCTTTCAGGGTGTGTTTATGAGTGACTCCAC CCAAGCTTGTAGCTGTTCTCTCCCATCTAACCTCAGGCAAGATCCTGGTGAAACAGCATG ACATGGCTTCTGGGGTGGAGGTTGGGGGTGGAGGTCCTGCTCCTAGAGATGAACTCTATC CAGCCCCTTAATTGGCAGGTGTATGTGCTGACAGTACTGAAAGCTTTCCTCTTTAACTGA TCCCACCCCACCCAAAAGTCAGCAGTGGCACTGGAGCTGTGGGCTTTGGGGAAGTCACT TAGCTCCTTAAGGTCTGTTTTTAGACCCTTCCAAGGAAGAGGCCAGAACGGACATTCTCT GCGATCTATATACATTGCCTGTATCCAGGAGGCTACACACCAGCAAACCGTGAAGGAGAA ACTTAAGCTCAATGTAACCCAGAGCCCACCATATAGTTTTATAGGTGCTCAACTTTCTAT

FIGURE 302

MGPVRLGILLFLFLAVHEAWAGMLKEEDDDTERLPSKCEVCKLLSTELQAELSRTGRSRE VLELGQVLDTGKRKRHVPYSVSETRLEEALENLCERILDYSVHAERKGSLRYAKGQSQTM ATLKGLVQKGVKVDLGIPLELWDEPSVEVTYLKKQCSTMLEEFEDIVGDWYFHHQEQPLQ NFLCEGHVLPAAETACLQETWTGKEITDGEEKTEGEEEQEEEEEEEEGGDKMTKTGSH PKLDREDL

Important features of the protein: Signal peptide: amino acids 1-21

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 106-110

N-myristoylation site: amino acids 115-121

Amidation site: amino acids 70-74

FIGURE 303

FIGURE 304

MMRCCRRRCCCRQPPHALRPLLLLPLVLLPPLAAAAAGPNRCDTIYQGFAECLIRLGDSM GRGGELETICRSWDFHACASQVLSGCPEEAAAVWESLQQEARQAPRPNNLHTLCGAPVH VRERCTGSETNQETLRATAPALPMAPAPPLLAAALALAYLLRPLA

Signal peptide:
Amino acids 1-35

Transmembrane domain: Amino acids 141-157

N-myristoylation site: Amino acids 127-133

Prokaryotic membrane lipoprotein lipid attachment site: Amino acids 77-88

FIGURE 305

FIGURE 306

MGWTMRLVTAALLLGLMMVVTGDEDENSPCAHEALLDEDTLFCQGLEVFYPELGNIGCKV VPDCNNYRQKITSWMEFIVKFFGAVDGATYILVMVDPDAPSRAEPRQRFWRHWLVTDIKG ADLKKGKIQGQELSAYQAPSPPAHSGFHRYQFFVYLQEGKVISLLPKENKTRGSWKMDRF LNRFHLGEPEASTOFMTONYQDSPTLQAPRGRASEPKHKTRQR

Important features of the protein: Signal peptide: amino acids 1-22

N-glycosylation site: amino acids 169-173

Tyrosine kinase phosphorylation site: amino acids 59-68

N-myristoylation sites: amino acids 54-60, 83-89, 130-136

Phosphatidylethanolamine signature: amino acids 113-157

FIGURE 307

 ${\tt AAGGAGCAGCCGCAAGCACCAAGTGAGAGGC} \underline{{\tt ATG}} \\ {\tt AAGTTACAGTGTTTTCCCTTTGGC}$ TCCTGGGTACAATACTGATATTGTGCTCAGTAGACAACCACGGTCTCAGGAGATGTCTGA TTTCCACAGACATGCACCATATAGAAGAGAGTTTCCAAGAAATCAAAAGAGCCATCCAAG CTAAGGACACCTTCCCAAATGTCACTATCCTGTCCACATTGGAGACTCTGCAGATCATTA AGCCCTTAGATGTGTGCTGCGTGACCAAGAACCTCCTGGCGTTCTACGTGGACAGGGTGT TCAAGGATCATCAGGAGCCAAACCCCAAAATCTTGAGAAAAATCAGCAGCATTGCCAACT CTTTCCTCTACATGCAGAAAACTCTGCGGCAATGTCAGGAACAGAGGCAGTGTCACTGCA GGCAGGAAGCCACCAATGCCACCAGAGTCATCCATGACAACTATGATCAGCTGGAGGTCC ACGCTGCTGCCATTAAATCCCTGGGAGAGCTCGACGTCTTTCTAGCCTGGATTAATAAGA ATCATGAAGTAATGTTCTCAGCTTGATGACAAGGAACCTGTATAGTGATCCAGGGATGAA CACCCCTGTGCGGTTTACTGTGGGAGACAGCCCACCTTGAAGGGGAAGGATGGGGAA GGCCCTTGCAGCTGAAAGTCCCACTGGCTGGCCTCAGGCTGTCTTATTCCGCTTGAAAA TAGGCAAAAAGTCTACTGTGGTATTTGTAATAAACTCTATCTGCTGAAAGGGCCTGCAGG CCATCCTGGGAGTAAAGGGCTGCCTTCCCATCTAATTTATTGTAAAGTCATATAGTCCAT GTCTGTGATGTGAGCCAAGTGATATCCTGTAGTACACATTGTACTGAGTGGTTTTTCTGA ATAAATTCCATATTTTACCTATGA

FIGURE 308

MKLQCVSLWILIGTILILCSVDNHGLRRCLISTDMHHIEESFOEIKRAIQAKDTFPNVTIL STLETLQIIKPLDVCCVTKNLLAFYVDRVFKDHQEPNPKIIRKISSIANSFLYMQKTLRQ CQEQRQCHCRQEATNATRVIHDNYDQLEVHAAAIKSLGELDVFLAWINKNHEVWFSA

Signal sequence: amino acids 1-18

N-glycosylation sites: amino acids 56-60, 135-139

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 102-106

N-myristoylation site: amino acids 24-30

Actinin-type actin-binding domain signature 1: amino acids 159-169

FIGURE 309

GTCGACCCACGCGTCCGAAGCTGCTGGAGCCACGATTCAGTCCCCTGGACTGTAGATAAA GACCCTTTCTTGCCAGGTGCTGAGACAACCACACTATGAGAGGCACTCCAGGAGACGCTG ATGGTGGAGGAAGGGCCGTCTATCAATCAATCACTGTTGCTGTTATCACATGCAAGTATC TGTGTTTGTATTGTGAGAAGGTTGGAGAACAGCCCACATTGCAGCTAAAAGAGCAGAAGA CTGGTAGGACCTCCACCCTTGAGTCTGTGGCCTTCCCGGACTGGTTCATTGCCTCCTCCA AGAGAGACCAGCCCATCATTCTGACTTCAGAACTTGGGAAGTCATACAACACTGCCTTTG AATTAAATATAAATGACTGAACTCAGCCTAGAGGTGGCAGCTTGGTCTTTGTCTTAAAGT TTCTGGTTCCCAATGTGTTTTCGTCTACATTTTCTTAGTGTCATTTTCACGCTGGTGCTG AGACAGGAGCAAGGCTGCTGTTATCATCTCATTTTATAATGAAGAAGAAGCAATTACTTC ATAGCAACTGAAGAACAGGATGTGGCCTCAGAAGCAGGAGAGCTGGGTGGTATAAGGCTG TCCTCTCAAGCTGGTGCTGTTGTAGGCCACAAGGCATCTGCATGAGTGACTTTAAGACTCA AAGACCAAACACTGAGCTTTCTTCTAGGGGTGGGTATGAAGATGCTTCAGAGCTCATGCG CGTTACCCACGATGGCATGACTAGCACAGAGCTGATCTCTGTTTCTGTTTTGCTTTATTC CCTCTTGGGATGATATCATCCAGTCTTTATATGTTGCCAATATACCTCATTGTGTGAAT TGCTTCAGAGCTCATGCGCGTTACCCACGATGGCATGACTAGCACAGAGCTGATCTCTGT TTCTGTTTTGCTTTATTCCCTCTTGGGATGATATCATCCAGTCTTTATATGTTGCCAATA TACCTCATTGTGTGTAATAGAACCTTCTTAGCATTAAGACCTTGTAAACAAAAATAATTC

FIGURE 310

MRGTPGDADGGGRAVYQSITVAVITCKYPEALEQGRGDPIYLGIQNPEMCLYCEKVGEQP TLQLKEQKIMDLYGQPEPVKPFLFYRAKTGRTSTLESVAFPDWFIASSKRDQPIILTSEL GKSYNTAFELNIND

Signal sequence: amino acids 1-17

N-myristoylation site: amino acids 10-16

Cell attachment sequence: amino acids 36-39

FIGURE 311

GCGAGGCTGCACCAGCGCCTGGCACCATGAGGACGCCTGGGCCTCTGCCCGTGCTGCTGC $\overline{\text{TGCTCCTGGCGGGAGCCCCGCCGCGCGCGCCACTCCCCCGACCTGCTACTCCCGCATGC}}$ GGGCCTGAGCCAGGAGATCACCCGCGACTTCAACCTCCTGCAGGTCTCGGAGCCCTCGG AGCCATGTGTGAGATACCTGCCCAGGCTGTACCTGGACATACACAATTACTGTGTGCTGG ACAAGCTGCGGGACTTTGTGGCCTCGCCCCCGTGTTGGAAAGTGGCCCAGGTAGATTCCT TGAAGGACAAAGCACGGAAGCTGTACACCATCATGAACTCGTTCTGCAGGAGAGATTTGG TATTCCTGTTGGATGACTGCAATGCCTTGGAATACCCAATCCCAGTGACTACGGTCCTGC GTCAGCTACCCAGACTTAATGGGCCAGAGCCATGACCCTCACAGGTCTTGTGTTAGTTGT ATCTGAAACTGTTATGTATCTCTCTACCTTCTGGAAAACAGGGCTGGTATTCCTACCCAG GAACCTCCTTTGAGCATAGAGTTAGCAACCATGCTTCTCATTCCCTTGACTCATGTCTTG CCAGGATGGTTAGATACACAGCATGTTGATTTGGTCACTAAAAAGAAGAAAAGGACTAAC AAGCTTCACTTTTATGAACAACTATTTTGAGAACATGCACAATAGTATGTTTTTATTACT GGTTTAATGGAGTAATGGTACTTTTATTCTTTCTTGATAGAAACCTGCTTACATTTAACC AAGCTTCTATTATGCCTTTTTCTAACACAGACTTTCTTCACTGTCTTTCATTTAAAAAAGA AATTAATGCTCTTAAGATATATTTTTACGTAGTGCTGACAGGACCCACTCTTTCATTGA AAGGTGATGAAAATCAAATAAAGAATCTCTTCACATGGA

FIGURE 312

MRTPGPLPVLLLLLAGAPAARPTPPTCYSRMRALSQEITRDFNLLQVSEPSEPCVRYLPR LYLDIHNYCVLDKLRDFVASPPCWKVAQVDSLKDKARKLYTIMNSFCRRDLVFLLDDCNA LEYPIPVTTVLPDRQR

Important features of the protein: Signal peptide: amino acids 1-19

Tyrosine kinase phosphorylation site: amino acids 60-69

N-myristoylation site: amino acids 16-22

FIGURE 313

GAGCGACGCTGTCTCTAGTCGCTGATCCCAAATGCACCGGCTCATCTTTGTCTACACTCT AATCTGCGCAAACTTTTGCAGCTGTCGGGACACTTCTGCAACCCCGCAGAGCGCATCCAT CAAAGCTTTGCGCAACGCCAACCTCAGGCGAGATGACTTGTACCGAAGAGATGAGACCAT CCAGGTGAAAGGAAACGGCTACGTGCAGAGTCCTAGATTCCCGAACAGCTACCCCAGGAA CCTGCTCCTGACATGGCGGCTTCACTCTCAGGAGAATACACGGATACAGCTAGTGTTTGA CAATCAGTTTGGATTAGAGGAAGCAGAAAATGATATCTGTAGGTATGATTTTGTGGAAGT TGAAGATATATCCGAAACCAGTACCATTATTAGAGGACGATGGTGTGGACACAAGGAAGT TCCTCCAAGGATAAAATCAAGAACGAACCAAATTAAAATCACATTCAAGTCCGATGACTA CTTTGTGGCTAAACCTGGATTCAAGATTTATTATTCTTTGCTGGAAGATTTCCAACCCGC AGCAGCTTCAGAGACCAACTGGGAATCTGTCACAAGCTCTATTTCAGGGGTATCCTATAA CTCTCCATCAGTAACGGATCCCACTCTGATTGCGGATGCTCTGGACAAAAAAATTGCAGA ATTTGATACAGTGGAAGATCTGCTCAAGTACTTCAATCCAGAGTCATGGCAAGAAGATCT TGAGAATATGTATCTGGACACCCCTCGGTATCGAGGCAGGTCATACCATGACCGGAAGTC AAAAGTTGACCTGGATAGGCTCAATGATGATGCCAAGCGTTACAGTTGCACTCCCAGGAA TTACTCGGTCAATATAAGAGAAGAGCTGAAGTTGGCCAATGTGGTCTTCTTTCCACGTTG CCTCCTCGTGCAGCGCTGTGGAGGAAATTGTGGCTGTGGAACTGTCAACTGGAGGTCCTG CACATGCAATTCAGGGAAAACCGTGAAAAAGTATCATGAGGTATTACAGTTTGAGCCTGG CCACATCAAGAGGAGGGGTAGAGCTAAGACCATGGCTCTAGTTGACATCCAGTTGGATCA CCATGAACGATGCGATTGTATCTGCAGCTCAAGACCACCTCGATAAGAGAATGTGCACAT CCTTACATTAAGCCTGAGAGAA

FIGURE 314

MHRLIFVYTLICANFCSCRDTSATPQSASIKALRNANLRRDDLYRRDETIQVKGNGYVQS PRFPNSYPRNLLLTWRLHSQENTRIQLVFDNOFGLEEAENDICRYDFVEVEDISETSTIT GRWCGHKEVPPRIKSRTNQIKITFKSDDYFVAKPGFKIYYSLLEDFQPAAASETNWESV TSSISGVSYNSPSVTDPTLIADALDKKIAEFDTVEDLLKYFNPESWQEDLENMYLDTPRY RGRSYHDRKSKVDLDRLNDDAKRYSCTPRNYSVNIREELKLANVVFFPRCLLVQRCGGNC GCGTVNWRSCTCNSGKTVKKYHEVLQFEPGHIKRRGRAKTMALVDIQLDHHERCDCICSS RPPR

Signal peptide: amino acids 1-18

N-glycosylation site: amino acids 270-274

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 262-266

Tyrosine kinase phosphorylation site: amino acids 256-265

N-myristoylation sites: amino acids 94-100, 186-192, 297-303, 298-304

TonB-dependent receptor proteins signature 1:
amino acids 1-56

FIGURE 315

CGGCTCGAGGCTCCCGCCAGGAGAAAGGAACATTCTGAGGGGAGTCTACACCCTGTGGAG CTCAAGATGGTCCTGAGTGGGGCGCTGTGCTTCCGAATGAAGGACTCGGCATTGAAGGTG GGTGAAGAGATCAGCGTGGTCCCCAATCGGTGGCTGGATGCCAGCCTGTCCCCCGTCATC CTGGGTGTCCAGGGTGGAAGCCAGTGCCTGTCATGTGGGGTGGGGCAGGAGCCGACTCTA ACACTAGAGCCAGTGAACATCATGGAGCTCTATCTTGGTGCCAAGGAATCCAAGAGCTTC ACCTTCTACCGGCGGACATGGGGCTCACCTCCAGCTTCGAGTCGGCTGCCTACCCGGGC TGGTTCCTGTGCACGGTGCCTGAAGCCGATCAGCCTGTCAGACTCACCCAGCTTCCCGAG AATGGTGGCTGGAATGCCCCCATCACAGACTTCTACTTCCAGCAGTGTGACTAGGGCAAC GTGCCCCCAGAACTCCCTGGGCAGAGCCAGCTCGGGTGAGGGGTGAGTGGAGGAGACCC ATGGCGGACAATCACTCTCTCTGCTCTCAGGACCCCCACGTCTGACTTAGTGGGCACCTG ACCACTTTCTCTCTCCTCCCAGTTTCGATAAATTCTGAGATTTGGAGCTCAGTCCACG GTCCTCCCCACTGGATGGTGCTACTGCTGTGGAACCTTGTAAAAACCATGTGGGGTAAA CTGCTTAATGGTAACTGACAAGTGTTACCCTGAGCCCCGCAGGCCAACCCATCCCCAGTT AGGGAGGTGGTCATAGAGTCAGGGATCTATGGCCCTTGGCCCAGCCCCACCCCCTTCCCT GCATGAGGAGGTGGTGATGTCAGAAGAAATGGCTCGAGCTCAGAAGATAAAAGATAAGTA GGGTATGCTGATCCTCTTTTAAAAACCCCAAGATACAATCAAAATCCCAGATGCTGGTCTC TATTCCCATGAAAAGTGCTCATGACATATTGAGAAGACCTACTTACAAAGTGGCATATA TTGCAATTTATTTAATTAAAAGATACCTATTTATATATTTCTTTATAGAAAAAAGTCTG GAAGAGTTTACTTCAATTGTAGCAATGTCAGGGTGGTGGCAGTATAGGTGATTTTTCTTT TAATTCTGTTAATTTATCTGTATTTCCTAATTTTTCTACAATGAAGATGAATTCCTTGTA TAAAAATAAGAAAAGAAATTAATCTTGAGGTAAGCAGAGCAGACATCATCTCTGATTGTC CTCAGCCTCCACTTCCCCAGAGTAAATTCAAATTGAATCGAGCTCTGCTGCTCTGGTTGG TTGTAGTAGTGATCAGGAAACAGATCTCAGCAAAGCCACTGAGGAGGAGGCTGTGCTGAG TTTGTGTGGCTGGATCTCTGGGTAAGGAACTTAAAGAACAAAAATCATCTGGTAATTCT $\tt TTCCTAGAAGGATCACAGCCCCTGGGATTCCAAGGCATTGGATCCAGTCTCTAAGAAGGC$ TGCTGTACTGGTTGAATTGTGTCCCCCTCAAATTCACATCCTTCTTGGAATCTCAGTCTG ATGAAGGTAGACCTAAATTCAATATGACTGGTTTCCTTGTATGAAAAGGAGGACACAG AGACAGAGGAGACGCGGGGAAGACTATGTAAAGATGAAGGCAGAGATCGGAGTTTTGCAG CCACAAGCTAAGAAACACCAAGGATTGTGGCAACCATCAGAAGCTTGGAAGAGGCAAAGA AGAATTCTTCCCTAGAGGCTTTAGAGGGATAACGGCTCTGCTGAAACCTTAATCTCAGAC TTCCAGCCTCCTGAACGAAGAAGAATAAATTTCGGCTGTTTTAAGCCACCAAGGATAAT TGGTTACAGCAGCTCTAGGAAACTAATACAGCTGCTAAAATGATCCCTGTCTCCTCGTGT TTACATTCTGTGTGTCCCCTCCCACAATGTACCAAAGTTGTCTTTTGTGACCAATAGAA TATGGCAGAAGTGATGCCATGCCACTTCCAAGATTAGGTTATAAAAGACACTGCAGCTTC GGGGGAAGCTAGCTGCCATGCTATGAGCAGGCCTATAAAGAGACTTACGTGGTAAAAAAT GAAGTCTCCTGCCCACAGCCACATTAGTGAACCTAGAAGCAGAGACTCTGTGAGATAATC GATGTTTGTTGTTTTAAGTTGCTCAGTTTTGGTCTAACTTGTTATGCAGCAATAGATAAA TAATATGCAGAGAAAGAG

FIGURE 316

MVLSGALCPEMKDSALKVLYLHNNQLLAGGLHAGKVIKGEEISVVPNRWLDASLSPVILG VQGGSQCLSCGVGQEPTLTLEPVNIMELYLGAKESKSFTFYRRDMGLTSSFESAAYPGWF LCTVPEADOPVRLTOLPENGGWNAPITDFYFOOCD

N-myristoylation sites: amino acids 29-34, 30-35, 60-65, 63-68, 73-78, 91-96, 106-111

Interleukin-1 signature: amino acids 111-131

Interleukin-1 proteins:
amino acids 8-29, 83-120, 95-134, 64-103

FIGURE 317

FIGURE 318

MELGLGGLSTLSHCPWPRRQPALWPTLAALALLSSVAEASLGSAPRSPAPREGPPPVLAS PAGHLPGGRTARWCSGRARRPPPQPSRPAPPPPAPPSALPRGGRAARAGGPGSRARAAGA RGCRLRSQLVPVRALGLGHRSDELVRFRFCSGSCRRARSPHDLSLASLLGAGALRPPPGS RPVSOPCCRPTRYEAVSFMDVNSTWRTVDRLSATACGCLG

signal sequence: Amino acids 1-39

N-glycosylation site: Amino acids 202-206

N-myristoylation sites: Amino acids 6-12;67-73;102-108;109-115;119-125

FIGURE 319

GTTGCTATGTTGCCCAGGCTGGTCTTGAAGTGCCTTGACCTCCTAAAGTGTTTGGAACCAC AGACGTGAGCCACTCCACCCAGCCTAAAACTTCATCTTCTTTGGATGAGATGAACACTTT TAACAAGAGAACAGGACTCTATATAAATCGCTGTGGGCTCACCACCTCTAAGGAGGAGCA CTGACTGAAGACAGAAAATTGATGAACTGAAGAAGACATGGTCCATTATGCCTTACAAA CTTACACAGTGCTTTGGGAATTCCAAAGTACTCAGTGGAGAGAGGTGTTTCAGGAGCCGT AGAGCCAGATCGTCATCATGTCTGCATTGTGGCTGCTGCTGGGCCTCCTTGCCCTGATGG ACTTGTCTGAAAGCAGCAACTGGGGATGCTATGGAAACATCCAAAGCCTGGACACCCCTG GAGCATCTTGTGGGATTGGAAGACGTCACGGCCTGAACTACTGTGGAGTTCGTGCTTCTG AAAGGCTGGCTGAAATAGACATGCCATACCTCCTGAAATATCAACCCATGATGCAAACCA TTGGCCAAAAGTACTGCATGGATCCTGCCGTGATCGCTGGTGTCTTGTCCAGGAAGTCTC CCGGTGACAAAATTCTGGTCAACATGGGCGATAGGACTAGCATGGTGCAGGACCCTGGCT CTCAAGCTCCCACATCCTGGATTAGTGAGTCTCAGGTTTCCCAGACAACTGAAGTTCTGA CTACTAGAATCAAAGAAATCCAGAGGAGGTTTCCAACCTGGACCCTGACCAGTACCTGA GAGGTGGACTCTGTGCCTACAGTGGGGGTGCTGGCTATGTCCGAAGCAGCCAGGACCTGA GCTGTGACTTCTGCAATGATGTCCTTGCACGAGCCAAGTACCTCAAGAGACATGGCTTCT AACATCTCAGATGAAACCCAAGACCATGATCACATATGCAGCCTCAAATGTTACACAGAT AAAACTAGCCAAGGGCACCTGTAACTGGGAATCTGAGTTTGACCTAAAAGTCATTAAAAT AACATGAATCCCATTAAAAAAAAAAAAAAA

FIGURE 320

MSALWLLLGLLALMDLSESSNWGCYGNIQSLDTPGASCGIGRRHGLNYCGVRASERLAEI DMPYLLKYQPMMQTIGQKYCMDPAVIAGVLSRKSPGDKILVNMGDRTSMVQDPGSQAPTS WISESQVSQTTEVLTTRIKEIQRRFPTWTPDQYLRGGLCAYSGGAGYVRSSQDLSCDFCN DVLARAKYLKRHGF

Important features of the protein: Signal peptide: amino acids 1-19

N-myristoylation sites: amino acids 23-29, 26-32, 35-41, 45-51, 50-56, 76-82, 156-162

Amidation site: amino acids 40-44

FIGURE 321

GCCTTATAAAGTAGCCTCTGCATCTGCCTGCCTCGGGCAGAGGAGGGCTACCCTGGGGCT GAGAGTTCACCTGTCTCAGGAACCACCTGAGCCCACAGATCCTGTGGGCAGCGGCCAGGG GTGCCTCCACGCCAGGCACCGTGGTCCGACTCAACAAGGCAGCATTGAGCTACGTGTCTG AAATTGGGAAAGCCCCTCTCCAGCGGGCCCTGCAGGTCACTGTCCCTCATTTCCTGGACT GGAGTGGAGAGGGGCTTCAGCCCACCAGGATCCGGATTCTGAATGTCCATGTGCCCCGCC ${\tt TCCACCTGAAATTCATTGCTGGTTTCGGAGTGCGCCTGCTGGCAGCAGCTAATTTTACTT}$ TATTCTCGGGCCACGCCAACGAGTTTGATGGCAGTAACAGCACCTCCCACGCGCTGCTGG TCCTGGTGCAGAAGCACATTAAAGCTGTCTTGAGTAACAAGCTGTGCCTGAGCATCTCCA ACCTGGTGCAGGGTGTCAATGTCCACCTGGGCACCTTAATTGGCCTCAACCCCGTGGGTC CTGAGTCCCAGATCCGCTATTCCATGGTCAGTGTGCCCACTGTCACCAGTGACTACATTT CCCTGGAAGTCAATGCTGTTCTCTTCCTGCTGGGCAACCCCATCATCCTGCCCACGGATG CCACCCTTTTGTGTTGCCAAGGCATGTGGGTACCGAGGGCTCCATGGCCACCGTGGGCC TCTCCCAGCAGCTGTTTGACTCTGCGCTCCTGCTGCTGCAGAAGGCCGGTGCCCTCAACC TGGACATCACAGGGCAGCTGAGGTCGGATGACAACCTGCTGAACACCTCTGCTCTGGGCC GGCTCATCCCGGAGGTGGCCCGCCAGTTTCCCGAGCCCATGCCTGTGGTGCTCAAGGTGC CCTTCGTGGAGGTCCTGGCCACAGCCTCCAACTCGGCTTTCCAGTCCCTCTTCTCCCTGG ATGTGGTAGTGAACTTGAGACTCCAGCTCTCTGTGTCCAAGGTGAAGCTTCAGGGGACCA CGTCTGTGCTGGGGGATGTCCAGCTCACGGTGGCCTCCTCCAACGTGGGCTTCATTGATA CAGATCAGGTGCGCACACTGATGGGCACCGTTTTTGAGAAGCCCCTGCTGGACCATCTCA ATGCTCTCTTGGCCATGGGAATTGCCCTCCCTGGTGTGGTCAACCTCCACTATGTTGCCC CTGAGATCTTTGTCTATGAGGGCTACGTGGTGATATCCAGTGGACTCTTCTACCAGAGCT CACCCAACACCACTCCCACCCCTGAGACTCGCCAGCTCGCTGCTCAGGCGAATTTCTCA TTTCAAGCCACTGGGGAAACTGAGGCAAAACCATACTTAGTCATCACCAACAAGCTGGAC CCCACCCAGGGGGGAGCAGACTGCTCCTCCAGGCTGTATAGACCTGCCCTCTTGCATTA AACAACTTCTCTTGAGCTGC

FIGURE 322

MAMASRLGILLALLIPVOGASTPOTVVRLNKAALSYVSEIGKAPLORALQVTVPHFLDWS
GEALQPTRIRILNVHVPRLHLKFIAGFGVRLLAANFTFKVFRAPEPLELTLPVELLADT
RVTQSSIRTPVVSISACSLFSGHANEFDGSNSTSHALLVLVQKHIKAVLSNKLCLSISNL
VQGVNVHLGTLIGLNPVGPESQIRYSMVSVPTVTSDYISLEVNAVLFILIGNPIILPTDAS
FFVLPRHVGTEGSMATVGLSQQLFDSALLLLQKAGALNLLDITGQLRSDDNLLNTSALGRL
IPEVARQFPEPMPVVLKVRLGATPVAMLHTNNATLRLQPFVEVLATASNSAFQSLFSLDV
VVNLRLQLSVSKVKLQGTTSVLGDVQLTVASSNVGFIDTDQVRTLMGTVFEKPLLDHLNA
LLAMGIALPGVVNLHYVAPEIFYVEGYVVISSKLFYGS

Important features of the protein: Signal peptide:
Amino acids 1-20

Transmembrane domain: Amino acids 217-236

N-glycosylation sites: Amino acids 96-100;151-155;293-297;332-336

N-myristoylation sites: Amino acids 8-14;149-155;189-195;249-255;252-258;283-289

LBP / BPI / CETP family proteins: Amino acids 22-50; 251-287

FIGURE 323

FIGURE 324

MKALMLLTLSVLLCWVSADIRCHSCYKVPVLGCVDRQSCRLEPGQQCLTTHAYLGKMWVF SNLRCGTPEEPCQEAFNQTNRKLGLTYNTTCCNKDNCNSAGPRPTPALGLVFLTSLAGLG LWLLH

Important features of the protein: Signal peptide: amino acids 1-18

N-glycosylation sites: amino acids 77-81, 88-92

N-myristoylation site: amino acids 84-90

Ly-6 / u-PAR domain protein signature: amino acids 85-98

FIGURE 325

ACGGCCGCAGCGCAGTGACGTAGGGTTGGCGCACGGATCCGTTGCGGCTGCAGCTCTG CAGTCGGGCCGTTCCTTCGCCGCCCCCGGGGTAGCGGTGTAGCTGCGCAGCGTCGCGCG CGCTACCGCACCCAGGTTCGGCCCGTAGGCGTCTGGCAGCCCGGCGCCATCTTCATCGAG TTTGTTTCTGCTGACCGCGGGCCCTGCCCTGGGCTGGAACGACCCTGACAGAATGTTGCT GCGGGATGTAAAAGCTCTTACCCTCCACTATGACCGCTATACCACCTCCCGCAGGCTGGA TCCCATCCCACACTTGAAATCTCTTTGCACCCACACCTGCTTGTTGATTCTTATACCCCAAA AGTCATACAGTGTCAGAACAAAGGCTGGGATGGGTATGATGTACAGTGGGAATGTAAGAC GGACTTAGATATTGCATACAAATTTGGAAAAACTGTGGTGAGCTGTGAAGGCTATGAGTC CTCTGAAGACCAGTATGTACTAAGAGGTTCTTGTGGCTTGGAGTATAATTTAGATTATAC AGAACTTGGCCTGCAGAAACTGAAGGAGTCTGGAAAGCAGCACGGCTTTGCCTCTTTCTC CGTGGTACTCCTTGGGATCGCCTTTGTAGTCTATAAGCTGTTCCTGAGTGACGGGCAGTA TTCTCCTCCACCGTACTCTGAGTATCCTCCATTTTCCCACCGTTACCAGAGATTCACCAA CTCAGCAGGACCTCCTCCCCCAGGCTTTAAGTCTGAGTTCACAGGACCACAGAATACTGG CCATGGTGCAACTTCTGGTTTTTGGCAGTGCTTTTACAGGACAACAAGGATATGAAAATTC CCCTGGCACGTGGAATAGGGCTTACTCACCCCTTCATGGAGGCTCGGGCAGCTATTCGGT ATGTTCAAACTCAGACACGAAAACCAGAACTGCATCAGGATATGGTGGTACCAGGAGACG ATAAAGTAGAAAGTTGGAGTCAAACACTGGATGCAGAAATTTTGGATTTTTCATCACTTT CTCTTTAGAAAAAAGTACTACCTGTTAACAATTGGGAAAAGGGGATATTCAAAAGTTCT GTGGTGTTATGTCCAGTGTAGCTTTTTGTATTCTATTATTTGAGGCTAAAAGTTGATGTG TGACAAAATACTTATGTGTTGTATGTCAGTGTAACATGCAGATGTATATTGCAGTTTTTG AAAGTGATCATTACTGTGGAATGCTAAAAATACATTAATTTCTAAAACCTGTGATGCCCT AAGAAGCATTAAGAATGAAGGTGTTGTACTAATAGAAACTAAGTACAGAAAATTTCAGTT TTAGGTGGTTGTAGCTGATGAGTTATTACCTCATAGAGACTATAATATTCTATTTGGTAT TATATTATTTGATGTTTTGCTGTTCTTCAAACATTTAAATCAAGCTTTGGACTAATTATGC TAATTTGTGAGTTCTGATCACTTTTGAGCTCTGAAGCTTTGAATCATTCAGTGGTGGAGA TGGCCTTCTGGTAACTGAATATTACCTTCTGTAGGAAAAGGTGGAAAATAAGCATCTAGA TCATAAGAGGTAAAGGTCAAATTTTTCAACAAAAGTCTTTTAATAACAAAAGCATGCAGT TCTCTGTGAAATCTCAAATATTGTTGTAATAGTCTGTTTCAATCTTAAAAAGAATCA

FIGURE 326

MAAACGPGAAGYCLLLGLHLFLLTAGPALGWNDPDRMLLRDVKALTLHYDRYTTSRRLDP IPQLKCVGGTAGCDSYTPKVIQCQNKGWDGYDVQWECKTDLDIAYKFGKTVVSCEGYESS EDQYVLRGSCGLEYNLDYTELGLQKLKESGKQHGFASFSDYYYKWSSADSCNMXGGLTTIV VLLGIAFVVYKLFLSDGQYSPPPYSEYPPFSHRYQRFTNSAGPPPPGFKSEFTGPQWTGH GATSGFGSAFTGQQGYENSGPGFWTGLGTGGILGYLFGSNRAATPFSDSWYYPSYPPSYP GTWNRAYSPLHGGSGSYSVCSNSDTKTRTASGYGGTRRA

Signal peptide:

Transmembrane domain: amino acids 171-190

N-glycosylation site: amino acids 172-176

Glycosaminoglycan attachment sites: amino acids 244-248, 259-263, 331-335

Tyrosine kinase phosphorylation site: amino acids 98-106

N-myristoylation sites: amino acids 68-74, 69-75, 131-137, 241-247, 247-253, 266-272, 270-276, 278-284, 312-318

FIGURE 327

FIGURE 328

MVSSWPARKASLLCVCAVLVLPWRTLGSPVILARRPGAWVPSWKGTSYTPQPHFPTNFYM PWENLLHVGCPLPLFQQCPVLLINLRPAPHTFPVQVPAVIPGSPMLLRPDGFLEAAGPWM

Signal peptide: amino acids 1-27

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 8-12

FIGURE 329

FIGURE 330

MQVSTAALAVLLCTMALCNQVLSAPLAADTPTACCFSYTSRQIPQNFIADYFETSSQCSK PSVIFLTKRGROVCADPSEEWVQKYVSDLELSA

Signal sequence: 1-23

Small cytokines (intercrine/chemokine) C-C subfamily signature:

1-35, 2-36, 10-44, 34-74, 50-90

Small cytokines (intecrine/chemokine): 24-89

FIGURE 331

 $\tt GGCACGAGGTGAGACTTTAAATGAA\underline{ATG} TCTCACAAGCTAGGTGATCCAGGTTTTGTGGT$ CTTTGCAACCCTTGTGGTCATTGTGGCCTTGATATTAATCTTCGTGGTGGTGGGTCCTCGCCA TCGACAGACAACATTCTTGTGTACATAACAATCTGCTCTGTAATCGGCGCGTTTTCAGT CCGCCATCCCCTGGCTTCGATTCTGCTGCTGAGCCTCATCGTCTGTGTGAGCACACAGAT TAATTACCTAAATAGGGCCCTGGATATATTCAACACTTCCATTGTGACTCCAATATATTA TGTATTCTTTACAACATCAGTTTTAACTTGTTCAGCTATTCTTTTTAAGGAGTGGCAAGA TATGCCTGTTGACGATGTCATTGGTACTTTGAGTGGCTTCTTTACAATCATTGTGGGGAT ATTCTTGCTTGCATGCCTTTAAAGACGTCAGCTTTAGTCTAGCAAGTCTGCCTGTGTCTTT TAATGAAGAAAGCTTAACCTGTGGAATCGAACACACACTGGTGAAAATGTCTCCCGAAG AAATGGAAATCTGACAGCTTTTTAAGAAAGGTGTAATTAAAGGTTAATCTGTGATTGTTA TGAAGTGAATTTGAATATCATCAGAATGTGTCTGAAAAAACATTGTCCTCAAATAATGTT $\tt CTTTAAAGGCAATCTTTTTAAAGATTTCACTAATTTGGACCAAGAAATTACTTTTCTTGT$ ATTTAAACAAACAATGGTAGCTCACTAAAATGACCTCAGCACATGACGATTTCTATTAAC ATTTTATTGTTGTAGAAGTATTTTACATTTTCATCCCTTCTCCAAAAGCCGAATGCACTA ATGACAGTTTTAAGTCTATGAAAATGCTTTATTTTTTCATTGGTGATGAAAGTCTGAAAT GTGCATTTGTCATCCCCACTCCATCAATCCCTGACCATGTAAGGCTTTTTTATTTTAAAA AAACAGAGTTATCCCAATACATTATCCTGTGATTTACCTTACCTACAAAAGTGGCTCCTG GAATGAAGGAACCTCTTTCTTACAAAACAAAAAAAGGGCAGAAATCACCCCAAGGAACG ATTTCTCAGGTTGAGATGATCACCGTGAATCCGGCTTCCTCTGAGCATTCGATGGCCTTA GTTACCCTAATCCCATGATGCCTGGAACCTTGATTACCGTTTTACATCAGCTCTTGTACT TTTCAGTATATTTCATAATGAGTTATATTGTCATTTAGACTTTGAACAGCTCTGGGAAA TAGAAGACTAGGGTTGTTTCTTAAATTTAGCTCATGTTATAATAAAAAGTTGAAATG

FIGURE 332

MSHKLGDPGFVVFATLVVIVALILIFVVGPRHGQTNILVYITICSVIGAFSVSCVKGLGI AIKELFAGRPVLRHPLAWILLLSLIVCVSTQINYLNRALDIFNTSIVTPIYYVFFTTSVL TCSAILFKEWQDMPVDDVIGTLSGFFTIIVGIFLLHAFKDVSFSLASLPVSFRKDEKAMN GNLSNMYEVLNNNEESLTCGIEQHTGENVSRRNGNLTAF

Signal sequence:

Transmembrane domain: 40-60, 70-90, 103-123, 139-159

N-glycosylation site: 103-106, 182-185, 208-211, 215-218

N-myristoylation site: 57-62, 140-145, 181-186, 214-219

FIGURE 333

GTGATGGCGGCTGGTGATGGGGACGTGAAGCTAGGCACCCTGGGGAGTGGCAGCGAGAGC AGCAACGACGCCGCAGCGAGAGTCCAGGCGACGCGGGAGCGGCAGCGGAAGGGGGAGGC TGGGCGGCGGCGTTGGCGCTTCTGACGGGGGGGGGGAAATGCTGCTGAACGTGGCG CTGGTGGCTCTGGTGCTGCTGGGGGCCTACCGGCTGTGGGTGCGCTGGGGGCGGGGGT CTGGGGGCCGGGGCGGGCGAGGAGAGCCCCGCCACCTCTCTGCCTCGCATGAAG AAGCGGGACTTCAGCTTGGAGCAGCTGCGCCAGTACGACGGCTCCCGCAACCCGCGCATC CTGCTCGCGGTCAATGGGAAAGTCTTCGACGTGACCAAAGGCAGCAAGTTCTACGGCCCG GCGGGTCCATATGGAATATTTGCTGGTAGGGATGCCTCCAGAGGACTGGCCACATTTTGC CTAGATAAAGATGCACTTAGAGATGAATATGATGATCTCTCAGATTTGAATGCAGTACAA ATGGAGAGTGTTCGAGAATGGGAAATGCAGTTTAAAGAAAAATATGATTATGTAGGCAGA CTCCTAAAACCAGGAGAAGAACCATCAGAATATACAGATGAAGAAGATACCAAGGATCAC AATAAACAGGATTGAACTTTGTAAACAACCAAAGTCAGGGGCCTTCAGAACTGCAATTCT TACTCCCTTTCACAGACTGTCCGGAGTCTTTGGGTTTGATTCACCTGCTGCGAAAAACAT TCAACAAATTGTGTACAAGATAAATTAATCTCACTATGAAGATTTGAATAACTAGACATT ATTTATGCTGCCAAACTCATTTGTTGCAGTTGTTTGTAATGTCTAGTGGGGGCTTCATCAT CCTGAAAAGAAGGAGACAGGGATTTTTTTAAAGAGCAAGAAAGTCACAATATTACTTCTT AAGACAACCAGATATGTATTTGCTACTCAAGTGTACAGATCTCCTCAAGAAACATCAAGG

FIGURE 334

MAAGDGDVKLGTLGSGSESSNDGGSESPGDAGAAAEGGGWAAAALALLTGGGEMLLNVAL VALVLLGAYRLWVRWGRRGLGAGAGAGEESPATSLPRMKKRDFSLEQLRQYDGSRNPRIL LAVNGKVFDVTKGSKFYGPAGPYGIFAGRDASRGLATFCLDKDALRDEYDDLSDLNAVQM ESVREWEMQFKEKYDYVGRLLKFGEEPSEYTDEEDTKDHNKQD

Signal sequence:

Transmembrane domain:

45-65

Tyrosine kinase phosphorylation site: 202-210

N-myristoylation site: 11-16, 16-21, 37-42, 38-43, 79-84, 81-86, 83-88, 144-149

Amidation site: 75-78

FIGURE 335

GACAGGCCGGGGTTACTGTGGCGACCACGAGAGCAGCTTTGGCGCTATGGAGGAGCCCGG GGCTACCCCTCAACCGTATTTGGGGCTGCTCCTGGAGGAGCTACGCAGGGTTGTGGCAGC ACTGCCTGAAGGTATGAGACCAGATTCTAATCTTTATGGTTTTCCATGGGAATTGGTGAT GGTTAGGAGTCGGCTTTATGTGGGACGAGAAAAAAGCTTGCTCTAATGCTTTCTGGACT AATTGAAGAAAAAAGTAAACTACTTGAAAAATTTAGCCTTGTTCAAAAAGAGTATGAAGG CTATGAAGTAGAGTCATCTTTAAAGGATGCCAGCTTTGAGAAGGAGGCAACAGAAGCACA AAGTTTGGAGGCAACCTGTGAAAAGCTGAACAGGTCCAATTCTGAACTTGAGGATGAAAT ACTCTGTCTAGAAAAAGAGTTAAAAGAAGAGAAATCCAAACATTCTGAACAAGATGAATT GATGGCGGATATTTCAAAAAGGATACAGTCTCTAGAAGATGAGTCAAAATCCCTCAAATC GATAGCAATAAAAGATGCTTTGAATGAAAATTCTCAACTTCAGGAAAGCCAGAAACAGCT TTTGCAAGAAGCTGAAGTATGGAAAGAACAAGTGAGTGAACTTAATAAACAGAAAGTAAC GACTCTGACTGAACGCTTGTTAAAGATGAAAGATTGGGCTGCTATGCTTGGAGAAGACAT AACGGATGATGATAACTTGGAATTAGAAATGAACAGTGAATCGGAAAATGGTGCTTACTT AGATAATCCTCCAAAAGGAGCTTTGAAGAAACTGATTCATGCTGCTAAGTTAAATGCTTC TTTAAAAACCTTAGAAGGAGAAAGAAACCAAATTTATATTCAGTTGTCTGAAGTTGATAA AACAAAGGAAGAGCTTACAGAGCATATTAAAAATCTTCAGACTCAACAAGCATCTTTGCA GTCAGAAAACACACATTTTGAAAATGAGAATCAGAAGCTTCAACAGAAACTTAAAGTAAT GACTGAATTATATCAAGAAAATGAAATGAAACTCCACAGGAAATTAACAGTAGAGGAAAA TTATCGGTTAGAGAAGAAGAAACTTTCTAAAGTAGATGAAAAGATCAGCCATGCCAC TGAAGAGCTGGAGACCTATAGAAAGCGAGCCAAAGATCTTGAAGAAGAATTGGAGAGAAC TATTCATTCTTATCAAGGGCAGATTATTTCCCATGAGAAAAAAGCACATGATAATTGGTT ACAAAAATTAACTGAAACAGAGCTTAAATTTGAACTTTTAGAAAAAGATCCTTATGCACT CGATGTTCCAAATACAGCATTTGGCAGAGGCTCACGAGGCCCAGGGAATCCTCTGGACCA TCAGATTACCAATGAAAGAGGAGAATCAAGCTGTGATAGGTTAACCGATCCTCATAGGGC TCCCTCTGACACTGGGTCTCTGTCACCTCCATGGGACCAGGACCGTAGGATGATGTTTCC TCCGCCAGGACAATCATATCCTGATTCAGCCCTTCCTCCACAAAGGCAAGACAGATTTTG TTCTAATTCTGGTAGACTGTCTGGACCAGCAGAACTCAGAAGTTTTAATATGCCTTCTTT GGATAAAATGGATGGGTCAATGCCTTCAGAAATGGAATCCAGTAGAAATGATACCAAAGA TGATCTTGGTAATTTAAATGTGCCTGATTCATCTCTCCCTGCTGAAAATGAAGCCACTGG CCCTGGCTTTGTTCCTCCACCTCTTGCTCCAATCAGAGGTCCATTGTTTCCAGTGGATGC AAGAGGCCCATTCTTGAGAAGAGGACCTCCTTTCCCCCCACCTCCTCCAGGAGCCATGTT TGGAGCTTCTCGAGATTATTTTCCACCAAGGGATTTCCCAGGTCCACCACCTGCTCCATT TGCAATGAGAAATGTCTATCCACCGAGGGGTTTTCCTCCTTACCTTCCCCCAAGACCTGG ATTTTTCCCCCCACCCCACATTCTGAAGGTAGAAGTGAGTTCCCCTCAGGTTTGATTCC ACCTTCAAATGAGCCTGCTACTGAACATCCAGAACCACAGCAAGAAACC<u>TGA</u>CAATATTT TTGCTCTCTCAAAAGTAATTTTGACTGATCTCATTTTCAGTTTAAGTAACTGCTGTTAC TTAAGTGATTACACTTTTGCTCAAATTGAAGCTTAATGGAATTATAATTCTCAGGATAGT ATTTTGTAAATAAAGATGATTTAAATATGAATCTTATGAGTAAATTATTTCAATTTTATT TTAGACGGTATAACTATTTCAATTTGATTAATCCACTATTATATAAACAATAGTGGGAGT TTTATATATGTAATCTTTCAGGTGGGGAGGCTTTAAATTCTGAAGTCTGTGTCTTTATGC TTATAGTTGATTTAAAGATTTGTTTGGCATTGATAATAATAAAATCAGTAGTTTTTCTAT

FIGURE 336

TGRGYCGDHESSFGAMEEPGATPQPYLGLLLEELRRVVAALPEGMRPDSNLYGFPWELVI CAAVVGFFFAVLFFLWRSFRSVRSRLYVGREKKLALMLSGLIEEKSKLLEKFSLVQKEYEG YEVESSLKDASFEKEATEAQSLEATCEKLNRSNSELEDELICLEKEBLKEEKSKHSEQDEL MADISKRIQSLEDESKSLKSQVAEAKNTFQIFQMNEERLKIAIKDALNENSQLQESQKQL LQEAEVWKEQVSELNKQKVTFEDSKVHAEQVLNDKESHIKTLTERLLKMKDWAAMLGEDI TDDDNLELEMNSESENGAYLDNPPKGALKKLIHAAKLNASLKTLEGERNQIYIQLEEVDK TKEELITEHIKNLQTQQASLQSENTHFENENQKLQKLKVMTELYQENEMKLHRKLTVEEN YRLEKEEKLSKVDEKISHATEELETYRKRAKDLEEELERTIHSYQGQIISHEKKAHDNWL AARNAERNLNDLRKENAHNRQKLTETELKFELLKEDPYALDVPNTAFGRGSRGPGNPLDH QITNERGESSCDRLTDPHRAPSDTGSLSPPWDQDRRWMFPPPGGSYPDSALPPQRQDRFC SNSGRLSGPAELRSFNMPSLDKMDGSMPSEMESSRNDTKDDLGNLNVPDSSLPAENEATG PGFVPPPLAPIRGFLFPVDARGPFLRRGPFFPPPPGAMFGASRDYFPPRFFGPPPPAF

Signal sequence:

Transmembrane domain: 54-74

N-glycosylation site: 150-153, 338-341, 636-639

cAMP- and cGMP-dependent protein kinase phosphorylation site: 413-416

Tyrosine kinase phosphorylation site: 414-421

N-myristoylation site: 466-417, 625-630, 697-702

Leucine zipper pattern: 142-163

FIGURE 337

GGACTGCGGTCTCGGGCAGCAATGGCCGAGAAGCGCGACACACGGGACTCCGAAGCCCAG TTGGTGGCGTTCTCATTCTTATTCACCGTTATAACTTTCCCAATCTCAATATGGATGTGC ATAAAGATTATAAAAGAGTATGAAAGAGCCATCATCTTTAGATTGGGTCGCATTTTACAA GGAGGAGCCAAAGGACCTGGTTTGTTTTTATTCTGCCATGCACTGACAGCTTCATCAAA GTGGACATGAGAACTATTTCATTTGATATTCCTCCTCAGGAGATCCTGACAAAGGATTCA GTGACAATTAGCGTGGATGGTGTGGTCTATTACCGCGTTCAGAATGCAACCCTGGCTGTG GCAAATATCACCAACGCTGACTCAGCAACCCGTCTTTTGGCACAAACTACTCTGAGGAAT GTTCTGGGCACCAAGAATCTTTCTCAGATCCTCTCTGACAGAAGAAGAAATTGCACACAAC ATGCAGTCTACTCTGGATGATGCCACTGATGCCTGGGGAATAAAGGTGGAGCGTGTGGAA ATTAAGGATGTGAAACTACCTGTGCAGCTCCAGAGAGCTATGGCTGCAGAAGCAGAAGCG CTGAAAGAAGCCTCCATGGTCATCACTGAATCTCCTGCAGCCCTTCAGCTCCGATACCTG CAGACACTGACCACCATTGCTGCTGAGAAAAACTCAACAATTGTCTTCCCTCTGCCCATA GATATGCTGCAAGGAATCATAGGGGCAAAACACAGCCATCTAGGC<u>TAG</u>TGTAGAGATGAG CGCTAGCCTTCCAAGCATGAAGTCGGGGACCAAATTAGCCTTTAACTCATAAAGAGAGGG TAGGGCTTTTCTTTTTCCATATGTCAATTGTGGTGTTCCCAGAATGTATAGCAGTTATAA AAATAGGTGAAAGAATTGTTAGCTTGTAAATACTGAGAGATTGGTGATTTATATAAGGTA ATCTGTTAGTCTTAAAATAGTTAAAAGTTTGTATTTTTAGATTATTATGTAGTTAGGTTAG ATCCCTCTTGTTTTGACTTCCACTGACTCATTCTGAACCCCCTAAGCACCCAGGCCACAG GCAAGAACCTGGGCTGTAACTGCCACCTGACACCGCTGACTGGCTAAATGCTTTGCAGAA AGTGATGACCTTACACCACAACCAGCTTCTCCAGGTCATATGTGCCTTACCTCCAGAAGT CTTTTTTTTTTTTTTTTTCTGAGATGGAGTTTCACTCTTGTTGCCCAGGCTGGAGTGCAA TAGCATGATCTCGGCTCACTGCAACCTCCGCCTCCTGGGTTCAAGAGATTCTCCTGCCTC AGCCTCCCAGTAGCTGGGATTACAGGCTCATGCCACCATGCCCAGCTAATTTTTGTATT ATTATTATTGTTTTTTAGTAGAGACGGGGTTTCACCATGTTGGCCAGGCTAGTCACGAAC TCCTAACCTCAGGTGATCCACCCACCTCTGCCTCCAAAGTGCTGGATTACAGGCTGAGCT AAATCCAGCCGTGTTTCAGAATAATCCTTACTTGAGAGTAGCCATTTTCTTGTGTACTTG TCAGAACTAGAGGAAATAGCCAAGACTAATGAAAAAACATTACTCTAACCCTTAAAAGACT TTTAAATTCACTACTAGAGTGGTCATTTTAAAAATACATCCATGTTTTAACTTATTTTGA GCCTTTCTTTTATGAGTAAATGATTCCTCCTTGTTCTGTCTTTCAAACCAGCTAAATATT TGTCACAAAAGTGACTTTTTCTCACTGTTGCCTATTTTCATATATCAGGTTTTAAATAG TTTTAATTTTTAATAAAATTTTTCTCTACGTTCTATATGCAATTGTTATATATCTATTT GAATAGCTGAAGGACTAAAATACTTTTTTAAGAGATAACTTCAGGAAACCATTATATTTT ACTATCTGCATGCTGTTAACTGTGGTACACTGTGAAATATGTTGATTACAAACCCATTCA TTACATAGTATAAGGAATTCACAGTATATTGACTATATAGTGTCTAATGACTGGGCAGAT ACTGTCAACTTACAATATCTATATAGAGAGGCTTTAAACTTACCTTACTCATTCTCTATG ATGTATGACTTGATGCTGAAAGAGGAAGCTGGTCAGCTCCTCATGGACAACAAATTCTTA GTCTATAATATTAGGAGACATCTCTAGTTTTGCAAATGTCTGTGAATCTGAGCAACCTGG ACTTCTGCTTACTGGCCAGAAAGCTGGCGGGTGACATTTGTAACATTTCCTCTTTGAGAC AGCTCTCTTTAGCTCAACCACTCTGTCCATCCAGCCAATGGATGTCCTTCCCTGTACCCA ATTCAAGCTTATTTTAGGGAAGCCTTGAAACTACCATGTATCTGGCTCTAGCTGAGTTAT TGAGGATTGAGCCAGTGCAACGTTAAACTCAGTGCACTTACATTTGATTTAAATGATGGT AAACCTTTTCTGAGACTTAGAGTAACAGTACTTTTGGTTCCTTGAGTTCTCCTGTCTCCA GATACCTAAATGACCTTGACTTTTCTGCCTTGTGAATTCGTAGTCCAATCAGCTGAAATT AAATCACTTGGGAGGGACGCATAGAAGGAGCTCTAGGAACACAGTGCCAGTGCAGAAGTT TCTCCAGGTGGCCTCCCTTTCCAACAATGTACATAATAAAGTGTATGCACTTTCACT

FIGURE 338

MAEKRDTRDSEAQRLPDSFKDSPSKGLGPCGWILVAFSFLFTVITFPISIWMCIKIIKEY ERAIIFRLGRILQGGAKGPGLFFILPCTDSFIKVDMRTISFDIPPQEILTKDSVTISVDG VVYYRVQNATLAVANITNADSATRLLAQTTLRNVLGTKNLSQILSDREEIAHNMQSTLDD ATDAWGIKVERVEIKDVKLPVQLQRAMAAEAEASREARAKVIAAEGEMNASRALKEASMV TTESPAALOLRYLOTLTTIAAEKNSTIVPPLPIDMLOGIIGAKHSHLG

Signal sequence:

Transmembrane domain: None

N-glycosylation site: 128-131, 135-138, 159-162, 229-232, 264-267

cAMP- and cGMP-dependent protein kinase phosphorylation site: $^{4-7}$

N-myristoylation site: 26-31, 278-283, 281-286

SPFH domain/Band 7 family: 39-230

FIGURE 339

TCTAGAGCCCTCTCCCAACATGGCGGCCTCAGCAAAAAAGAAGAATAAGAAGGGGAAGAC TATCTCCCTAACAGACTTTCTGGCTGAGGATGGGGGTACTGGTGGAGGAAGCACCTATGT TTCCAAACCAGTCAGCTGGGCTGATGAAACGGATGACCTGGAAGGAGATGTTTCGACCAC TTGGCACAGTAACGATGACGATGTGTATAGGGCGCCTCCAATTGACCGTTCCATCCTTCC CACTGCTCCACGGGCTGCTCGGGAACCCAATATCGACCGGAGCCGTCTTCCCAAATCGCC ACCCTACACTGCTTTTCTAGGAAACCTACCCTATGATGTTACAGAAGAGTCAATTAAGGA ÅTTCTTTCGAGGATTAAATATCAGTGCAGTGCGTTTACCACGTGAACCCAGCAATCCAGA GAGGTTGAAAGGTTTTGGTTATGCTGAATTTGAGGACCTGGATTCCCTGCTCAGTGCCCT GAGTCTCAATGAAGAGTCTCTAGGTAACAGGAGAATTCGAGTGGACGTTGCTGATCAAGC ACAGGATAAAGACAGGGATGATCGTTCTTTTGGCCGTGATAGAAATCGGGATTCTGACAA AACAGATACAGACTGGAGGGCTCGTCCTGCTACAGACAGCTTTGATGACTACCCACCTAG AAGAGGTGATGATAGCTTTGGAGACAAGTATCGAGATCGTTATGATTCAGACCGGTATCG GGATGGGTATCGGGATGGGTATCGGGATGGCCCACGCCGGGATATGGATCGATATGGTGG CCGGGATCGCTATGATGACCGAGGCAGCAGAGACTATGATAGAGGCTATGATTCCCGGAT AGGCAGTGGCAGAAGAGCATTTGGCAGTGGGTATCGCAGGGATGATGACTACAGAGGAGG CGGGGACCGCTATGAAGACCGATATGACAGACGGGATGATCGGTCGTGGAGCTCCAGAGA TGATTACTCTCGGGATGATTATAGGCGTGATGATAGAGGTCCCCCCAAAGACCCAAACT GAATCTAAAGCCTCGGAGTACTCCTGAAGAAGATGATTCCTCTGCTAGTACCTCCCAGTC CACTCGAGCTGCTTCTATCTTTGGAGGGGCAAAGCCTGTTGACACAGCTGCTAGAGAAAG AGAAGTAGAAGAACGGCTACAGAAGGAACAAGAGAAGTTGCAGCGTCAGTGGAATGAGCC AAAACTAGAACGACGGCCTCGGGAGAGACACCCAAGCTGGCGAAGTGAAGAAACTCAGGA ACGGGAACGGTCGAGGACAGGAAGTGAGTCATCACAAACTGGGACCTCCACCACATCTAG CAGAAATGCACGAAGGAGAGAGTGAGAAGTCTCTAGAAAATGAAACACTCAATAAGGA GGAAGATTGCCACTCTCCAACTTCTAAACCTCCCAAACCTGATCAGCCCCTAAAGGTAAT GCCAGCCCTCCACCAAAGGAGAATGCTTGGGTGAAGCGAAGTTCTAACCCTCCTGCTCG ATCTCAGAGCTCAGACACAGAGCAGCAGTCCCCTACAAGTGGTGGGGGAAAAGTAGCTCC AGCTCAACCATCTGAGGAAGGACCAGGAAGGAAAGATGAAAATAAAGTAGATGGGATGAA TGCCCCAAAAGGCCAAACTGGGAACTCTAGCCGTGGTCCAGGAGACGGAGGGAACAGAGA CCACTGGAAGGAGTCAGATAGGAAAGATGGCAAAAAGGATCAAGACTCCAGATCTGCACC $\tt TGCTCTCTGTTGATGGTGAAGATGAAAATGAGGGAGAAGATTATGCCGAA\underline{TAG}ACCTC$ TACATCCTGTGCTTTTCTCCTAGTTTCTCTCCACCCTGGAACATTCGAGAGCAAATCAAA ACCTCTATCCAGACAAGACAAAATAAAACTCAACATCTCCTGAAGACCTTTCTTACCTTT TTTTAAAAACAAAAXTGAAATTATTTTGCATGCTGCTGCAGCCTTTAAAGTATTGAAGT AACTGGAGAATTGCCAATACAGCCAGAGAGAAAGGGACTACAGCTTTTTAGAGGAAAAGT TGTGGTGCGTTATGTCACCATGCAGTTGCCAGTGTGATTAGTGCCTAGGGGTCTCATTTA GGTAGAAGGAAAAGTGTGAGATTTCTACCTTTTAGTTTTCATCCTATTGTGGCATATATG AATTCTCAAACATTATCTGAATAAATTTTCCACTCTTGGAAAGGTAGATTTAGCCTCAAG TTGTTCTAGTCTCCAGGAGGCTGCCAGCCCCTCCTCTTATTTAATTCTGAGTTTTGGGGG CCAGCCTAGAGGGAATTCCTTTTTTTTTTTTTTAACCCCCCAGGGGGGTAGTTGGGAGT TTTGAAAAGAAGCTTTTGGGAAGTGATGAGTCATTTTGCACCAGGTAATAGGGGAAAATT GTGTGACCTCCAGCAAACACATGAATGGTTATTTCCTGGAGCCGGAAGCACTTGGGGGTC GTGGTAATTCCCAGTGTTTTCTGTGTCCTAGTTTTACCCTTTCTAAACACTGTCCTTTTT GAAAGTTTTGAATATATCCACATTCTATTGAAACCTTGAAACTAAAAATTTAGACTCTTA TCGTCATCTTAAGTTCTTCATGCTACTCTTAACCTCCCAAAAAGCAGTATCTAAGTCACA TACATGATGTCTTGGGCATTTTCTGAGCCATGGAGAACTCTGAAAGGAAGAATCGCTGCT TTTCTCAAGCAAATCGGTTTCTTGATGTCTTTTGGTTCTCCTTGCCTGCTCCTGATGCTT

TAGGGACAGGACAGTTAAATTGGGAGCCTTTCTTACAACCTTGATGGGATTTTTCCCCC CAAGTTTCCTTCTCCACTGAAATGCCACACTAATGCTTGTTGGATTCATGAGGTGGCCAG ACCAATGTGTTGTTTGTTGTTTTTTTTTTTAAGCTTCCCTTGAGAGAATAAATGGTA ATGGAGAGAATCATTTAACAAGGTCCTGGTTTCTCTTGCAACACAGTAGCTAAACTTGCC TGCTTTTATATGCATTTTTGTAGGGATCAGCTTGGTAGACAGTATTAGCGGAGAAACACC TTGATCTTGGTTTGCAAGCCCTTCTCCCATCAGTCCTAGATTAGGCCCTGTTCAGCCATG CAGGGGTGTTGGTTTATGCGTGCTGCAGCAGTGGGCATAATGAATATAATTTACCCAGTG GACAAAGGTGTGTACCAAGTGAATTTAAATAATTGGTGTGGATTGGCCAGTAGCTAAGAA AACAAACTATTGATTGTAGATAATGAAAAGCTAGGGTTTGCCCTCTTCATGTCTACTCTC CTTCCAAATAGTTATATCCAAAACTGTTTTTCCCTCTCCCCTACCTTGTCCCCCCTATTA AAATAGAAACAGGGATTGATTAATGTCCCGCTCCTGAATACATGTAAAATTTGTACAAAA ATATCTTCTATGAAAATGATTTGTAATCTGTAGACTTATTACCTGGGAGATGTCTTGATG AGTTAAAAAAAAAAAAAAACTCTAGAGTCGAGGAATTC

FIGURE 340

MAASAKKNKKGKTISLTDFLAEDGGTGGGSTYVSKPVSWADETDLLEGDVSTTWHSNDD
DVYRAPPIDRS ILPTAPRAREENIDRSRLPKSPPYTAFLGNLPYDVTEESIKEFFRGLN
ISAVRLPREPSNPERLKGFGYABFEDLDSLLSALSLMEESLGNRRIRVDVADQAQDKDRD
DRSFGRDRNRDSDKTDTDWRARPATDSFDDYPPRRGDDSFGDKYRDRYDSDRYRDGYRDG
RYDGPRRDMDRYGGRDRYDDRGSRDYDRGYDSRIGSGRRAFGSGYRRDDDYRGGGDRYED
RYDRRDDRSWSSRDDYSRDDYRRDDRGPPQRPKLNLKPRSTPEEDDSSASTSQSTRAASI
FGGAKPVDTAAREREVEERLQKEQEKLQRQWNEPKLERRPRERHPSWRSEETQEREKSFT
SESSQTGTSTTSSRNARRESEKSLENETLINKEEDCHSPTSKPPKPDQPLKVMPAPPPK
ENAWVKRSSNPPARSQSDTEQQSPTSGGGKVAPAQPSEEGPGRKDENKVDGMNAPKGQT
GNSSRGPGDGGRNRHWKESDRKDGKKDQDSRSAPEPKKPEENPASKFSSASKYAALSVDG

```
Signal Sequence:
None
Transmembrane domain:
None
N-glycosylation site:
120-123, 448-451, 542-545
Glycosaminoglycan attachment site:
507-510
cAMP- and cGMP-dependent protein kinase phosphorylation
site:
439-442, 486-489
Tyrosine kinase phosphorylation site:
225-233, 264-270
N-myristoylation site:
25-30, 26-31, 28-33, 118-123, 421-426, 428-433, 538-543
Amidation site:
276-279, 522-525, 563-566
Cell attachment sequence:
215-217
Eukaryotic putative RNA-binding region RNP-1 signature:
137-144
RNA recognition motif:
```

98-168

FIGURE 341

GCGTGGACACCACCTCAGCCCACTGAGCAGGAGTCACAGCACGAAGACCAAGCGCAAAGC GACCCTGCCTCCATCCTGACTGCTCCTCCTAAGAGAGATGGCACCGGCCAGAGCAGGA GCCAAGCCCAAGGGCATGACCTCATCACAGTGGTTTAAAATTCAGCACATGCAGCCCAGC CCTCAAGCATGCAACTCAGCCATGAAAAACATTAACAAGCACACAAAACGGTGCAAAGAC CTCAACACCTTCCTGCACGAGCCTTTCTCCAGTGTGGCCGCCACCTGCCAGACCCCCAAA ATAGCCTGCAAGAATGGCGATAAAAACTGCCACCAGAGCCACGGGCCCGTGTCCCTGACC ATGTGTAAGCTCACCTCAGGGAAGTATCCGAACTGCAGGTACAAAGAGAAGCGACAGAAC AAGTCTTACGTAGTGGCCTGTAAGCCTCCCCAGAAAAAGGACTCTCAGCAATTCCACCTG GTTCCTGTACACTTGGACAGAGTCCTTTAGGTTTCCAGACTGGCTTGCTCTTTGGCTGAC CTTCAATTCCCTCTCCAGGACTCCGCACCCCCTACACCCAGAGCATTCTCTTCCCC TCATCTCTTGGGGCTGTTCCTGGTTCAGCCTCTGCTGGGAGGCTGAAGCTGACACTCTGG TCCCCAAGAAACAGCAAGCTCAGGTCTGTGGGTTCCCTGGTCTATGCCATTGCACATGTC

FIGURE 342

MAPARAGFCPLLLLLLGLWVAEIPVSAKPKGMTSSQWFKIQHMQPSPQACNSAMKNINK HTKRCKDLNTFLHEPFSSVAATCQTPFLIACKNGDKNCHQSHGPVSLTMCKLTSGKYPNCR YKEKRONKSYVVACKPPQKKDSQQFHLVPVHLDRVL

Important features of the protein

Signal peptide: 1-22

Transmembrane domain:

N-glycosylation site:

cAMP- and cGMP-dependent protein kinase phosphorylation site: 139-143

N-myristoylation site: 18-24, 32-38

Pancreatic ribonuclease family signature: 65-72

Pancreatic ribonuclease family proteins: 49-93

FIGURE 343

GCATTTGCCACTGGTTGCAGATCAGGCGGACGAGGGCCGGGAAGGCAGAGCCATGTGGC CTGTGAGAGCCCCAGAGCAGGGGTCCCTGACGGTTCAATGCCACTATAAGCAAGGATGGG AGACCTACATTAAGTGGTGGTGCCGAGGGGTGCGCTGGGATACATGCAAGATCCTCATTG AAACCAGAGGGTCGGAGCAAGGAGAGAGAGTGACCGTGTGTCCATCAAGGACAATCAGA AAGACCGCACGTTCACTGTGACCATGGAGGGGCTCAGGCGAGATGACGCAGATGTTTACT GGTGTGGGATTGAAAGAAGAGGACCTGACCTTGGGACTCAAGTGAAAGTGATCGTTGACC CAGAGGGAGCGCTTCCACAACAGCAAGCTCACCTACCAACAGCAATATGGCAGTGTTCA TCGGCTCCCACAAGAGGAACCACTACATGCTCCTGGTATTTGTGAAGGTGCCCATCTTGC TCATCTTGGTCACTGCCATCCTCTGGTTGAAGGGGTCTCAGAGGGTCCCTGAGGAGCCAG GGGAACAGCCTATCTACATGAACTTCTCCGAACCTCTGACTAAAGACATGGCCACT**TAG**A GAGATGGATCTGCAGAGCCTTCCTGCCCTGGCCACGTTTCCAGAAGAGACTCGGGCTGTG GAAGGAACATCTACGAGTCCTCGGGATGCAGTGACTGAGATAGGGGCCCTGGGCCTCCGC CCTGGCCTTGGAGCTGGTGGGCACCTCCCTGTTCTGCACAGCTCAGGGACTTAGCCAGGT CCTCTCCTGAGCCACCATCACCTCCTGGGGTGCCAGCACCTGTTCTCTTGGTCAGGAGCT GTAGAGATGGAGCTCAAGCACTGGACGACTCTGTCCCCACTGCTGGAATAACTCGGGCAC AGAGCATGGGACCAAAGTACAGAAAGAGGTTGGGGGAGACCCCCCCAGCCCTAGACTTCC ATCATTCCGGAGACCAACTCAACACCGTCTTTGCCTGAGAACCTGATATATCCGTGTTTT TAAATTTTTTTTTTTCTAGCAAAGTTGGGTTTTAATGACTTATGTTCATAGGAAACCTCT CTGATCCCACACACAGGGGGGTGATTCTGGGATGAGTTCCTGGTTCTAGGGCATGAGGG GCTGGATGGACCCTGTCCCCAGGGAGGACATGGCTCTGAGTCCACAGGGCTGAGGAGGCA TAGCTCCCAAGCTCCCTGCCTATTCCCCCACCTCCGAGGGGCTGCAGCTTGGGAGCCTC CTCAGCATGACAGCTTGGGTCTCCTCCCCAAAAGAGCCTGTCAGGCCTCAAGAACCACCT ATGTTGAAATCATGTTACTAATGAAAACTGTCCTAGGGAAGTGGTTCTGTCTCCTCACAG GCTTCACCCACGGCGATGAGGCCCTTGAATGTGGTCACTTTGTGCTGTATGGTTGAGGGA CCCTCACACCAAAGGGACCTTCCCATGTGAGATGTGCTCCCGCCCCCACCTGCCCACAAG CAAACACACCACACGTTCGGCATGTTGCCCTTTGAACACCCATGAGGACGCCTCCAAC CTGCTCTTGGTTCTAATAGGGAGTACTGACTGTCAGCAGTGGATAAAGGAGAGGGGACCC TCTGGTCCCTAGCATGGCACCCAGAGCCTCCCCTCTTCTTGTCCTTCAGCCAAAGAGAAA CTTTCTCTGACTTTGAACTGAATTTAGGTCTCTGGCCAATGATGGGCCTGAAAATTCCAT AATGGCCAGAGAGAGAGTTCGAGCCCGGCTAAGATCCCCTGAGTCATTCTGTGAGGGAC CAAGACCCACAGTCCACCAGCCCCAGGGCCCTACCTCCTGGAATGCTTTCCTGGATCCAG AAGCATGGTGCTTCACCAGCTGGACTCAGGGGGCGAGGGGACATGGGCGCTTGTCAACGTG ATGTCATTCTTTTCCCACCGTTTCTTCCTGTTGATATTCAATGAATCCGTCAATCTCTCT

FIGURE 344

MWLPPALLLLSLSGCFSIQGPESVRAPEQGSLTVQCHYKQGWETYIKWWCRGVRWDTCKI LIETRGSEQGEKSDRVSIKDNQKDRTFTVTMEGLRRDDADVYWCGIERRGPDLGTQVKVI VDPEGAASTTASSPTNSNMAVFIGSHKRNHYMLLVFVKVPILLILVTAILWLKGSQRVPE EPGEOPIYMNFSEPLTKDMAT

Important features of the protein: Signal peptide: Amino acids1-17

Transmembrane domain:

N-glycosylation site: Amino acids 190-194

Tyrosine kinase phosphorylation site: Amino acids 95-103

N-myristoylation sites: Amino acids 66-72;125-131

Prokaryotic membrane lipoprotein lipid attachment site: Amino acids5-16

FIGURE 345

CTGAGCTCCCGGGCTCCGCCAGCGCGCTGGCGGGGCGCCCCATTGCACACTCTGGGGGCG CCGCAGTGTTCGTGGGATGGGGCAGCGGGCTGCAGCTGGCGGCCGGAATCCGCGCGCAGC CCGGGTGCAAGTTCTCTCCTGTTGCCCTGAGTGCCCACTCCCAGGCCCTCTGTATGAGTG ACACTTCAGTCTGCCATGGAACCTGGCCCTGCTCTGGCCTGGCTCCTGCTCCTGAGCCTG CTGGCGGATTGTCTGAAAGCTGCTCAGTCCCGAGACTTCACAGTGAAAGACATTATCTAC CTCCATCCTTCAACCACACCATATCCTGGTGGATTTAAATGTTTCACCTGTGAAAAGGCA GCAGACAATTATGAGTGCAACCGATGGGCTCCAGACATCTACTGCCCTCGAGAGACCAGA TACTGCTACACTCAGCACACAATGGAAGTCACAGGAAACAGTATCTCAGTCACCAAACGC TGTGTCCCACTGGAAGAGTGCTTATCCACTGGCTGCAGAGACTCCGAGCATGAAGGCCAC AAGGTCTGCACTTCTTGTTGTGAAGGAAATATCTGTAACTTGCCACTGCCCCGAAATGAA ACTGATGCCACATTTGCCACGACGTCACCTATAAATCAGACAAATGGGCACCCACGCTGT ATCTCACTCATACTCTCCTCCTTCTCGCTTGTGGTTAGGGCTCATGTTATAGTGGCTCAGT GCTCCATGTGTTAATAGCGATCCATGGGGATCTCGATGGTCCACAGACCTGCATGAGTC ATTGGCCTGACAGTAATTACACATGTGAGACACACACTCTTGGAGGTCATCACAGCCAA GCATTGCCACTTACCATGAGGAATAAATGTTGCTTCATTGTAGCCATTTTGAGTCTAACC CATTCCAAGAAGTAGTTCTGCATTTATCGAGATCTGGGGTTCTTAATTTGGAAGAATACA TGCATGAGATGCAGTAGGTCCTGAGACTGTAAGATATTAGGAGTATGTTATAGGGGCCATG TATAGATGTGGGCTTTTCAGGAGAAAAGTAACCATTGGTTTAAATATAATCATGAGTTCA TTTGTAGCTTTAGAATTTTAAAACATTGACTCCAAACTGAATGGACTATTTCCTTGGAAA TTCTGACTGAGTCCCTGGAAGAGTAGTAATTCCAACAATTCCAGCCATTTGTTCAATTAA TTTTCCCAACATTCTTCTCCCAGTGCTGGGAATCACATTTCCTCTGTTCTGTGCAGAAGA CAAAAAGGCAATCATAAAAGTTTGTTATATTTGTGGGGGTGCCTGGAGGAGGATTTTCCT CAACTTAATGGAGCCACTGTCCATAAAGTGGCTGTTATCCCTTCATAATAATTGGTGAGAT CAGCCTTCTCCTTGACTTGGCACCTAATTATGCTTCATGAGATCCTAGATTCCACCTGAG TCAATTGTGTCCAGAGCCCCAAACCAGGATGGAGTTGTTTTCCCCAGATATGGGGTTCTA TTCAGCCATAGATAATCTAGACAGAGGATTTCAGAATGAAAGGAAAAATGTGTGGAGATT GTTCATACTGCCAAAGAGCTCCCACTTCCAAATCCCCAGTGACTTTATGGAGAAGATTCT GCATTAAATTGTCTTTCGAATGATGGGGAAGCAAGGCATAATATGCGATGATGAGGAGAA TTTTAGATATTGCTTTTGAAGTAGATGGTAAAATTTTTGTCATCCTTCTTGTATTTTTTG AAGGCATAACTAGAAACTAAAATATATTCTAAAAAATTCATTATTCTGAACAAAGTGATC AAATTAGAATACATATTTTTCAACAGTGGTAGAGCTTTTAATATATGTTTATTGAAAGTT ATCTATAATACTTGCACCAGTGTTGAAAAAAGTTAACATGTAGGCAAGAGCAATATGTTT GTCTCAAGGATTTTTCCATGGTTTCCTCAGTGATGGTGTCCTGGAATTATTCAGGTGGTG ACCATCACTGGTCTAAGTTTGTGTGCAGGGTTTTCAGACGTGTTTTTGTGAAACTTGGTA GAACCATGGCTAATAAAGAGGACAGTGTTGTCAGGGTCCATCTGCCCTCCATAGAAAAAT GTCTCTGGCTCATAAAATGAGACTCCCTCAGGGACTAAATATGAACTGACAGCAGTAACT CTGATACAGAATAATCTAAATTGCATCAAATGGCCTTAATTCAGAGTTTGTTAGGCTTAT CAGTATGTTGCTTTTAATTGGGGTGGGAAAGTAGAGGGAGAGAAAGCAAGACATTTATTA AGCACCTCGTATGTGCCAGGCACTATGCTAAGCACTTTACATAAGTTAGGATTAATCCCT GCAAGAATCCTATAAAGAATGTTACTAGCATTTACACTTCCCAAATGAAGGTACCAAAGC TCAAACGCAATGTTGTGAAGCTGTTTCCTTCAGATTTAGGTTATGTGGGATGATGTGGGA TTGAAGAGGAAAGAAGGTGGGATTATCCCCCTAGGAAGACTTTCAGGCCTGACTTCATA GGAATTCATCCATCTTATCATGTGGAGTTTATCTCACCCTGCTGTTGCAGGATGCTATTT

FIGURE 346

MEPGPALAWLLLLSLLADCLKAAQSRDFTVKDIIYLHPSTTPYPGGFKCFTCEKAADNYE CNRWAPDIYCPRETRYCYTQHTMEVTGNSISVTKRCVPLEECLSTGCRDSEHEGHKVCTS CCEGNICNLPLPRNETDATFATTSPINQTNGHPRCMSVIVSCLWLWLGLML

Important features of the protein:

Signal peptide: 1-22

Transmembrane domain: None

N-glycosylation site: 134-138, 147-151

N-myristoylation site: 45-51, 87-93, 106-112, 124-130

Ly-6 / u-PAR domain protein: 115-128

FIGURE 347

GATCAAGCGCCTTCCTTTCCCTTCCTCTCCCTACTTGGCCCTTTGCCCTAAGCCAAGACCT GGCCATCAGCCTGCCAGGGGCCTGCAGAGCCAGCTGCACTTTTTCAGGTATGGGGGA $\tt GGGCCAGGCACC{\color{red}{\textbf{ATG}}} {\tt AAGCCAGTGTGGGTCGCCACCCTTCTGTGGATGCTACTGCTGGTG$ CCCAGGCTGGGGGCCCCGGAAGGGGTCCCCAGAAGAGGCCTCCTTCTACTATGGAACC TTCCCTCTTGGCTTCTCCTGGGGCGTGGGCAGTTCTGCCTACCAGACGGAGGGCGCCTGG GTGCTTGGGAATGAGACGCAGATGTAGCCTGTGACGGCTACTACAAGGTCCAGGAGGAC ATCATTCTGCTGAGGGAACTGCACGTCAACCACTACCGATTCTCCCTGTCTTGGCCCCGG CTCCTGCCCACAGGCATCCGAGCCGAGCAGGTGAACAAGAAGGGAATCGAATTCTACAGT GATCTTATCGATGCCCTTCTGAGCAGCAACATCACTCCCATCGTGACCTTGCACCACTGG GATCTGCCACAGCTGCTCCAGGTCAAATACGGTGGGTGGCAGAATGTGAGCATGGCCAAC TACTTCAGAGACTACGCCAACCTGTGCTTTGAGGCCTTTGGGGACCGTGTGAAGCACTGG ATCACGTTCAGTGATCCTCGGGCAATGGCAGAAAAAGGCTATGAGACGGGCCACCATGCG CCGGGCCTGAAGCTCCGCGGCACCGGCCTGTACAAGGCAGCACACCACATCATTAAGGCC GGAATTTCACTGAACTGTGACTGGGGGGAACCTGTGGACATTAGTAACCCCAAGGACCTA GAGGCTGCCGAGAGATACCTACAGTTCTGTCTGGGCTGGTTTGCCAACCCCATTTATGCC GAGATGTCGAGGTTACCGGTGTTCTCACTCCAGGAGAGAGCTACATTAAAGGCACATCC GATTTCTTGGGATTAGGTCATTTTACTACTCGGTACATCACGGAAAGGAACTACCCCTCC CGCCAGGGGCCCAGCTACCAGAACGATCGTGACTTGATAGAGCTGGTTGACCCAAACTGG CCAGATCTGGGGTCTAAATGGCTATATTCTGTGCCATGGGGATTTAGGAGGCTCCTTAAC TTTGCTCAGACTCAATACGGTGATCCTCCCATATATGTGATGGAAAATGGAGCATCTCAA AAATTCCACTGTACTCAATTATGTGATGAGTGGAGAATTCAATACCTTAAAGGATACATA AATGAAATGCTAAAAGCTATAAAAGATGGTGCTAATATAAAGGGGGTATACTTCCTGGTCT GAATTTAACGACAGAAATAAGCCTCGCTATCCAAAGGCTTCAGTTCAATATTACAAGAAG ATTATCATTGCCAATGGGTTTCCCAATCCAAGAGAGGGTGGAAAGTTGGTACCTCAAAGCT TTGGAAACTTGCTCTATCAACAATCAGATGCTTGCTGCAGAGCCTTTGCTAAGTCACATG CAAATGGTTACGGAGATCGTGGTACCCACTGTCTGCTCCCTCTGTGTCCTCATCACTGCT GTTCTACTAATGCTCCTCCTGAGGAGGCAGAGCTGAGACAGGATTATCAATTTTGGAGCT TCATAAGAGAATCTTCAGGATCTTCCTCCCTTTTCTGCTTTGAGGGTTTCCATACATTGC TGTTTTCAGGTTCTACAATAATTACCTTTTTTTCTCTTTTCTCTTTTTGGCTTGTGCTGGG ATTTAAGAATTAGAAAATAAAAATAAGCAGAAATTA

FIGURE 348

MKPVWVATLLWMLLLVPRLGAARKGSPEEASFYYGTFPLGFSWGVGSSAYQTEGAWDQDG
KGPSIWDVFTHSGKGKVLGNETADVACDGYYKVQEDIILLRELHVNHYRFSLSWPRLLPT
GIRAGVMKKGIEFYSDLIDALLSSNTFPLTVTHHWDLDPQLLQVKYGGWQNVSMANVFRD
YANLCFEAFGDRVKHWITFSDPRAMAEKGYETGHHAPGLKLRGTGLYKAAHHIIKAHAKT
WHSYNTTWRSKQQGLVGISLNCDWGEPVDISNPKDLEAAERYLQFCLGWFANPIYAGDYP
QVMKDYIGRKSAEQGLEMSRLPVFSLQEKSYIKGTSDFLGLGHFTTRYITERNYBSRQGP
SYQNDRDLIELVDPNWPDLGSKWLYSVPWGFRRLLWFAQTQYGDPPIYVMENGASQKFHC
TQLCDEWRIQYLKGYINEMLKAIKDGANIKGYTSWSLLDKFEWEKGYSDRYGFYYVEFND
RNKPRYPKASVQYYKKIILANGFPNPREVESWYLKALETCSINNQMLAAEPLLSHWQMVT

Important features:
Signal peptide:
amino acids 1-21

Transmembrane domain: amino acids 541-558

N-glycosylation sites: amino acids 80-84,171-175,245-249

Glycosaminoglycan attachment site: amino acids 72-76

cAMP- and cGMP-dependent protein kinase phosphorylation sites: amino acids 23-27,564-568

Tyrosine kinase phosphorylation sites: amino acids 203-211,347-355,460-468,507-514

N-myristoylation sites: amino acids 44-50.79-85,167-173,225-231,257-263,315-321

Amidation site: amino acids 307-311

Glycosyl hydrolases family 1 active site: amino acids 407-416

Glycosyl hydrolases family 1 N-terminal signature: amino acids 41-56

Motif name Glycosyl hydrolases family: amino acids 37-67

FIGURE 349

CGCAAAGCCGCCTCGGGGCGCTCATGGCGGGACGCCTCCTGGGAAAGGCTTTAGCCGCG GTGTCTCTCTCTGGCCTTGGCCTCTGTGACTATCAGGTCCTCGCGCTGCCGCGCATC CAGGCGTTCAGAAACTCGTTTTCATCTTCTTGGTTTCATCTTAATACCAACGTCATGTCT GGTTCTAATGGTTCCAAAGAAAATTCTCACAATAAGGCTCGGACGTCTCCTTACCCAGGT GACTATAAGCCTGTGGAATACACTGCAGTCTCTGTCTTGGCTGGACCCAGGTGGGCAGAT CCTCAGATCAGTGAAAGTAATTTTTCTCCCAAGTTTAACGAAAAGGATGGGCATGTTGAG AGAAAGAGCAAGAATGGCCTGTATGAGATTGAAAATGGAAGACCGAGAAATCCTGCAGGA CGGACTGGACTGGTGGGCCGGGGGCTTTTGGGGCCGATGGGGCCCAAATCACGCTGCAGAT CCCATTATAACCAGATGGAAAAGGGATAGCAGTGGAAATAAAATCATGCATCCTGTTTCT GGGAAGCATATCTTACAATTTGTTGCAATAAAAAGGAAAGACTGTGGAGAATGGGCAATC CCAGGGGGGATGGTGGATCCAGGAGAGAGATTAGTGCCACACTGAAAAGAGAATTTGGT GAGGAAGCTCTCAACTCCTTACAGAAAACCAGTGCTGAGAAGAGAGAAATAGAGGAAAAAG TTGCACAAACTCTTCAGCCAAGACCACCTAGTGATATATAAGGGATATGTTGATGATCCT CGAAACACTGATAATGCATGGATGGAGACAGAAGCTGTGAACTACCATGACGAAACAGGT GAGATAATGGATAATCTTATGCTAGAAGCTGGAGATGATGCTGGAAAAGTGAAATGGGTG GACATCAATGATAAACTGAAGCTTTATGCCAGTCACTCTCAATTCATCAAACTTGTGGCT GAGAAACGAGATGCACACTGGAGCGAGGACTCTGAAGCTGACTGCCATGCGTTG<u>TAG</u>CTG ATGGTCTCCGTGTAAGCCAAAGGCCCACAGAGGAGCATATACTGAAAAGAAGGCAGTATC ACAGAATTTATACTATAAAAAGGGCAGGGTAGGCCACTTGGCCTATTTACTTTCAAAACA ATTTGCATTTAGAGTGTTTCGCATCAGAATAACATGAGTAAGATGAACTGGAACACAAAA GCATGGCTTAAATTAAATTTAAACAACTAATGCTCTTTGAAGAATCATAATCAGAATAAA GATAAATTCTTGATCAGCTATA

FIGURE 350

MAGRILGKALAAVSLSLALASVTIRSSRCRGIQAFRNSFSSSWFHLNTNVMSGSNGSKEN SHNKARTSBYBGKVERSQVPMEKVGRLVEWQDYKPVEYTAVSVLAGPRWADPQISESN SPKFNEKDGHVERKSKNGLYEI ENGRPRNPAGRTGLUGRGGLIGRWGPNHAADPIITRWKR DSSGNKIMHPVSGKHILQFVAIKRKDCGEWAIPGGMVDPGEKISATLKREFGEEALNSLQ KTSAEKREIEEKLHKLFSQDHLVIYKGYVDDPRNTDNAWMETEAVNYHDETGEIMDNLML EAGDDAGKVKWVDINDKLKLYASHSOFIKLVAEKRDAHWSEDSEADCHAL

Important features of the protein:

Signal peptide:

1-20

Transmembrane domain:

N-glycosylation site:

55-59

cAMP- and cGMP-dependent protein kinase phosphorylation site: 179-183

N-myristoylation site: 53-59, 56-62

mutT domain signature: 215-235

FIGURE 351

 $\tt CCTCTGTCTGTGCTCCCATCCCAGGGGGGTATAGGTGGAGCCTCCAGAGCCCATGGACAGG$ GCATGCTGGGGCTGGGCCAGCCCAGCGGTGTCTCTAAGGCACCCCTGGGATCCCCACTG AGCTGGCCTACTTCAGACAGCCAGGGCCCACCCCTCTGGCCCCCTTAGTGTCCAGCTCGT GGCCCTTGGCATTTCCACAAGACGCCAAGATGGAGATTCCCATGGGGACCCAGGGCTGC TTCTCAAAGAGCCTCCTGCTCTCAGCCTCAATCCTGGTCCTCTGGATGCTCCAAGGCTCC CAGGCAGCTCTCTACATCCAGAAGATTCCAGAGCAGCCTCAAAAGAACCAGGACCTTCTC CTGTCAGTCCAGGGTGTCCCAGACACCTTCCAGGACTTCAACTGGTACCTGGGGGAGGAG ACGTACGGAGGCACGAGGCTATTTACCTACATCCCTGGGATACAACGGCCTCAGAGGGAT GGCAGTGCCATGGGACAGCGAGACATCGTGGGCTTCCCCAATGGTTCCATGCTGCTGCGC CGCGCCCAGCCTACAGACAGTGGCACCTACCAAGTAGCCATTACCATCAACTCTGAATGG ACTATGAAGGCCAAGACTGAGGTCCAGGTAGCTGAAAAGAATAAGGAGCTGCCCAGTACA CACCTGCCCACCAACGCTGGGATCCTGGCGGCCACCATCATTGGATCTCTTGCTGCCGG GCCCTTCTCATCAGCTGCATTGCCTATCTCCTGGTGACAAGGAACTGGAGGGCCCAGAGC CACAGACTGCCTGCTCCGAGGGGCCAGGGATCTCTGTCCATCTTGTGCTCGGCTGTATCC CCAGTGCCTTCAGTGACGCCCAGCACATGGATGGCGACCACAGAGAAGCCAGAATTGGGC CCTGCTCATGATGCTGGTGACAACAACATCTATGAAGTGATGCCCTCTCCAGTCCTCCTG GTGTCCCCCATCAGTGACACAAGGTCCATAAACCCAGCCCGGCCCCTGCCCACACCCCCA CACCTGCAGGCGGAGCCAGAAACCACCAGTACCAGCAGGACCTGCTAAACCCCGACCCT GAAGACAAGGCCCCAGCCCTCCTCTGGGAGCCTCACACCTGAGACCAGCAGGACAAGGCC ATTGGGGGCTGTGGGGCCGATGAGGTGGACTCAGCCAAAGACTCAGCAGCACATGGGGCA GGTGTCCTGGCAGGGGGACAGGACTGTAACAGGCCCAGGTCCTTGTGCAGCCCCTGAA TGCACGCCCGCCTTCGGTCTGTTCCTTCAAGCAAGCTGGCCTGGGCCATGTGCCTGTGAA AGGCAGGCTCTGGCCCCTTTCCATGCCAAAGTCCCCCAAGATCTGGATATCTGGGGACAA ATGCCCTACCCCAACTCCACTAGTGACCCTCAGAGTCTTCTCCCCTTAGGACAAGGCAGA CACCCCACCATGCGGGCCTCAGGTGGCAGAGAGGCCCAGCCTCACAGGCCTGTGGCCCCA AGCCCCTCTCAGAACCTGCTGCCAGCTGCTGGTCTTGGCCCCCACCCTGAATCTTACTGA GCA

FIGURE 352

MEIPMGTQGCFSKSLLLSASILVLWMLQGSQAALYIQKIPEQPQKNQDLLLSVQGVPDTF QDFNWYLGEETYGGTRLFTYIPGIQRPQRDGSAMGQRDIVGFPNGSMLLRRAQPTDSGTY QVAITINSEWTMKAKTEVQVAEKNKELPSTHLPTNAGILAATIIGSLAAGALLISCIAYL LVTRNWRGQSHRLPAPRGQGSLSILCSAVSPVPSVTPSTWMATTEKPELGPAHDAGDNNI YEVMPSPVLLVSPISDTRSINPARPLPTPPHLQAEPENHQYQQDLLNPDPAPYCQLVPTS

Important features of the protein: Signal peptide: Amino acids 1-32

Transmembrane domain: Amino acids 159-178

N-glycosylation site: Amino acids 104-108

N-myristoylation sites: Amino acids6-12;29-35;55-61;91-97;157-163;165-171

FIGURE 353

CTTCAGAACAGGTTCTCCTTCCCCAGTCACCAGTTGCTCGAGTTAGAATTGTCTGCAATG GCCGCCTGCAGAAATCTGTGAGCTCTTTCCTTATGGGGACCCTGGCCACCAGCTGCCTC CTTCTCTTGGCCCTCTTGGTACAGGGAGGAGCAGCTGCGCCCATCAGCTCCCACTGCAGG CTTGACAAGTCCAACTTCCAGCAGCCCTATATCACCAACCGCACCTTCATGCTGGCTAAG GAGGCTAGCTTGGCTGATAACAACACAGACGTTCGTCTCATTGGGGAGAAACTGTTCCAC GGAGTCAGTATGAGTGAGCGCTGCTATCTGATGAAGCAGGTGCTGAACTTCACCCTTGAA GA AGTGCTGTTCCCTCA ATCTGATAGGTTCCAGCCTTATATGCAGGAGGTGGTGCCCTTC CTGGCCAGGCTCAGCAACAGGCTAAGCACATGTCATATTGAAGGTGATGACCTGCATATC CAGAGGAATGTGCAAAAGCTGAAGGACACAGTGAAAAAGCTTGGAGAGAGTGGAGAGATC AAAGCAATTGGAGAACTGGATTTGCTGTTTATGTCTCTGAGAAATGCCTGCATT**TGA**CCA GAGCAAAGCTGAAAAATGAATAACTAACCCCCTTTCCCTGCTAGAAATAACAATTAGATG CCCCAAAGCGATTTTTTTTAACCAAAAGGAAGATGGGAAGCCAAACTCCATCATGATGGG TGGATTCCAAATGAACCCCTGCGTTAGTTACAAAGGAAACCAATGCCACTTTTGTTTATA AGACCAGAAGGTAGACTTTCTAAGCATAGATATTTATTGATAACATTTCATTGTAACTGG TGTTCTATACACAGAAAACAATTTATTTTTTAAATAATTGTCTTTTTCCATAAAAAAAGAT TACTTTCCATTCCTTTAGGGGAAAAAACCCCTAAATAGCTTCATGTTTCCATAATCAGTA ATTAATATGGATTTATTTATAGAAACATCATTCGATATTGCTACTTGAGTGTAAGGCTAA TATTCATATTTTCACAATATTATACACCTATAACATCTTTATTCACCTCAATAACA CTTGGATATCCC

FIGURE 354

MAALQKSVSSFLMGTLATSCLLLLALLVQGGAAAPISSHCRLDKSNFQQPYITNRTFMLA KEASLADNNTDVRLIGEKLFHGVSMSERCYLMKQVLNFTLEEVLFFQSDRFQPYMQEVVP FLARLSNRLSTCHIEGDDLHIORNVOKLKDTVKKLGESGEIKAIGBLDLLFMSLRNACI

Important features of the protein: Signal peptide: amino acids 1-33

N-glycosylation sites: amino acids 54-58, 68-72, 97-101

N-myristoylation sites: amino acids 14-20, 82-88

Prokaryotic membrane lipoprotein lipid attachment site: amino acids 10-21

FIGURE 355

TGGCCTACTGGAAAAAAAAAAAAAAAAAAAAAAAGTCACCGGGCCCGCGGTGGCCACAA CCGGGCCAGTCGGATCTCAGCCACGGACGGCGTTTCTCGGACCTCAAAGTGTGCGGGGA CGAAGAGTGCAGCATGTTAATGTACCGTGGGAAAGCTCTTGAAGACTTCACGGGCCCTGA TTGTCGTTTTGTGAATTTTAAAAAAGGTGACGATGTATATGTCTACTACAAACTGGCAGG GGGA TCCCTTGA ACTTTGGGCTGGA AGTGTTGA ACACAGTTTTGGATATTTTCCAAAAGA TTTGATCAAGGTACTTCATAAATACACGGAAGAGAGCTACATATTCCAGCAGATGAGAC TTCTCAGCACAGAGAAATCTCCTGAGGAGTCTCGGGGGGCGTGAACTTGACCCTGTGCC TGAGCCCGAGGCATTCAGAGCTGATTCAGAGGATGGAGAAGGTGCTTTCTCAGAGAGCAC CGAGGGGCTGCAGGGACAGCCCTCAGCTCAGGAGAGCCACCCTCACACCAGCGGTCCTGC GGCTAACGCTCAGGGAGTGCAGTCTTCGTTGGACACTTTTGAAGAAATTCTGCACGATAA ATTGAAAGTGCCGGGAAGCGAAAGCAGAACTGGCAATAGTTCTCCTGCCTCGGTGGAGCG

FIGURE 356

MAAAPGLLFWLFVLGALWWVPGQSDLSHGRRFSDLKVCGDEECSMLMYRGKALEDFTGPD CRFVNFKKGDDVYVYYKLAGGSLELMAGSVEHSFGYFFKDLIKVLHKYTEEELHIFADET DFVCFEGGRDFNSYNVEELLGSLELEDSVPEESKKAEEVSGHREKSPEESRGRELDPVP EPEAFRADSEDGEGAFSESTEGLQGQPSAQESHPHTSGPAANAQGVQSSLDTFEEILHDK LKVPGSESRTGNSSPASVEREKTDAYKVLKTEMSQRGSGQCVIHYSKGFRWHQNLSLFYK DCF

Important features of the protein: Signal peptide: amino acids 1-22

N-glycosylation site: amino acids 294-298

cAMP- and cGMP-dependent protein kinase phosphorylation site: amino acids 30-34

Tyrosine kinase phosphorylation site: amino acids 67-76

N-myristoylation sites: amino acids 205-211, 225-231, 277-283

Amidation site: amino acids 28-32

FIGURE 357

ACGCGCCGGCAGCTGTCCACCGATCCCGGCCACCCCCCGGCCACCCCCACCCCGCGA GCCCATGGAGGCTCCGGGACCCCGCGCCTTGCGGACTGCGCTCTGTGGCGGCTGTTGCTG CCTCCTCTATGTGCCCAGCTGGCTGTGGCTGGTAAAGGAGCTCGAGGCTTTGGGAGGGG AGCCCTGATCCGCCTGAATATCTGGCCGGCGGTCCAAGGGGCCTGCAAACAGCTGGAGGT CTGTGAGCACTGCGTGGAGGGAGACAGAGCGCGCAATCTCTCCAGCTGCATGTGGGAGCA GAAGACAGTCACAACAGGGAGCCCCCAGTCCCTGAGGCCCACAGCCCTGGATTTGACGG GCTGCACTTCCTCAAGGCCAAGGACACCTACCAGACGCTGTGAGTACCTGGCCAGCA GCAAGTACCTGAGTCCCAGCTCACCTCCTGGTTCCTGCCCCACCGTTCCCCTTCAGTACC CAGGGTGCTGTCTCCCATGGGCAAGCCCTCAGGACGGTGACAGCGTGCTCCATGTGAG CCACACCCCTTTTGTCTCCTCCAGTTGGGGTGTTTCCTTTGTCAGATGTTGGCTGGGACC AGGACTCAGCCTGGGCCAGTCTAGGAGCCCAGCTGAGCCCTCCTGTGTCTTTTCCCTTCA TGCTGCCAGCAGGAAGAACCAGTAGGTGCCAGCCCAGGCAAGCCTGTGGCCCGCGTT TCTGTGGCTGTGGGCAGGAGCTGGGCCTTGTGTCTAGTTGGGTTTTGCTCTGAGAAGGGG AGCTGTGCCTGAGGCCCTCTGTGTGCCGTGTGTGCTGTGGGGCGGGTCGCCACAGCCTGT GAGGCAGTGTCACCTTCCTGAGTGTCCTCTTTGGCCTCTGCAGAATCTGACCCCTTTGGG $\tt CCTGGACTCCATCCTGAGGGGAAAGGAGGATGCAGAGGGTGGCCTCTGGGCACCCTTGTG$ GGTAAGCGGGGGGGGGGGGGGAAAAACTCTGGCCGCCAGTTTTTGGCTCCTGCGGGCA CCAAGCAGGCTCAGTGTCTGATGCCTGACATCTCCTCCTGTCCTGGGCCTGGAACCTGCA GCTGAGAAAATCCCTCAACCACCTCGTCTCCTCCATCGCCCCTGCTGGGCCCCCAGCCT GACAGTGGGTTGTATGCCTGCCTCTTTCCACCAACTGGCCTGGGCACTGCCCCCAAATAA AAAAAAAAAAAAAACCA

FIGURE 358

MEAPGPRALRTALCGGCCCLLLCAQLAVAGKGARGFGRGALIRLNIWPAVQGACKQLEVC EHCVEGDRARNLSSCMWEQCRPEEPGHCVAQSEVVKEGCSIYNRSEACPAAHHHPTYEPK TVTTGSPPVPEAHSPGFDGASFIGGVVLVLSLQAVAFFVLHFLKAKDSTYQTL

Important features of the protein:

Signal peptide: 1-29

Transmembrane domain: 141-160

N-glycosylation site: 71-75, 103-107

Tyrosine kinase phosphorylation site: 164-171

N-myristoylation site:

FIGURE 359

FIGURE 360

MARILLLFLPGLVAVCAVHGIFMDRLASKKLCADDECVYTISLASAQEDYNAPDCRFINV KKGQQIYVYSKLVKENGAGEFWAGSVYGDGQDEMGVVGYFPRNLVKEQRVYQEATKEVPT TDIDFFCE

Important features of the protein:

Signal peptide: 1-14

Transmembrane domain:

N-myristoylation site: 84-90

FIGURE 361

GGCACGAGCCACCACTTACAACCACACAGCCTATCCAGAAACATGAAGATAAGAAATGCT TGTGCTGTCCTTATTGAAGTACTCCTGTTTATACTTGAAGGAGTTACAGGAGCTCGAAAA ATTTCAACTTTCTCAGGCCCTGGCTCATGGCCGTGCAATCCCAAGTGTGATGGCAGAACT TACAACCCCTCAGAGGAGTGTTGTGTTCATGACACCATCCTGCCCTTTAAGCGGATTAAC $\tt CTCTGTGGCCCTAGCTGCACCTACAGGCCCTGCTTTGAGCTCTGCTGTCCTGAGTCCTAT$ TCATCCCCTATCTCCAGGAACTGTAAAAGCAACAAGATTTTTCATGGAGAAGATATTGAA ${\tt GACAACCAACTTTCTCTTAGGAAAAAAGTGGTGACCAGCCT} {\tt TGA} {\tt GAGTCTGCTTTCTTC}$ GCAGTGATATATGTGTCTCATTACAATGCTCCTTTGGATATTGTTTTCCTAAGCATGTGT TGGAATGTTCCCCCATAACTTTCTAAAATTATCCTATTTCAATGCAACTAAAGATAAATG ATCCACAGGACCTTTTCAAGATTTTAGAAGCAGCAAACTATGGCTGAGAGAAAAGACTCT CTGACCAGGCAAATTGTTCTGCAGTATTCTCCGGGCGTGTAGCTCCCCTGAGTAGTCGCC AGGCTGGTCTTGGCTTTGTAATAATACAGCTGCCTTTGAGTCCTCCCTACCCTGTTAGTA ACCCCTTGCCTGCACTGTTGTCCTTACAACCGAAATAAACTGATTAGTTG

FIGURE 362

MKIRNACAVLIEVLLFILEGVTGARKISTFSGPGSWPCNPKCDGRTYNPSEECCVHDTIL PFKRINLCGPSCTYRPCFELCCPESYSPKKKFIVKLKVHGERSHCSSSPISRNCKSNKIF HGEDIEDNOLSLRKKSGDQP

Important features of the protein:

Signal peptide: 1-23

Transmembrane domain:

Glycosaminoglycan attachment site: 31-35

N-myristoylation site: 20-26, 34-40

FIGURE 363

ACACTGGCCAAACAAAAACGAAAGCACTCCGTGCTGGAAGTAGGAGGAGAGTCAGGACTC CCAGGACAGAGAGTGCACAAACTACCCAGCACAGCCCCCTCCGCCCCCTCTGGAGGCTGA AGAGGGATTCCAGCCCCTGCCACCCACAGACACGGGCTGACTGGGGTGTCTGCCCCCCTT GGGGGGGGCACCACGGCCTCAGGCCTGGGTGCCACCTGGCACCTAGAAG<u>ATG</u>CCTGT GCCCTGGTTCTTGCTGTCCTTGGCACTGGGCCGAAGCCCAGTGGTCCTTTCTCTGGAGAG GCTTGTGGGGCCTCAGGACGCTACCCACTGCTCTCCGGGCCTCTCCTGCCGCCTCTGGGA CAGTGACATACTCTGCCTGCCTGGGGACATCGTGCCTGCTCCGGGCCCCGTGCTGGCGCC TACGCACCTGCAGACAGAGCTGGTGCTGAGGTGCCAGAAGGAGACCGACTGTGACCTCTG TCTGCGTGTGGCTGTCCACTTGGCCGTGCATGGGCACTGGGAAGAGCCTGAAGATGAGGA AAAGTTTGGAGGAGCAGCTGACTCAGGGGTGGAGGAGCCTAGGAATGCCTCTCTCCAGGC CCAAGTCGTGCTCTCCTTCCAGGCCTACCCTACTGCCCGCTGCGTCCTGCTGGAGGTGCA AGTGCCTGCTGCCCTTGTGCAGTTTGGTCAGTCTGTGGGCTCTGTGGTATATGACTGCTT CGAGGCTGCCCTAGGGAGTGAGGTACGAATCTGGTCCTATACTCAGCCCAGGTACGAGAA TGACAACGTGCATCTGGTTCTGAATGTCTCTGAGGAGCAGCACTTCGGCCTCTCCCTGTA CTGGAATCAGGTCCAGGGCCCCCAAAACCCCGGTGGCACAAAAACCTGACTGGACCGCA GATCATTACCTTGAACCACACAGACCTGGTTCCCTGCCTCTGTATTCAGGTGTGGCCTCT GGAACCTGACTCCGTTAGGACGAACATCTGCCCCTTCAGGGAGGACCCCCGCGCACACCA ACCGTGCTCGCTGCCCGCAGAAGCGGCACTGTGCTGGCGGGCTCCGGGTGGGGACCCCTG CCAGCCACTGGTCCCACCGCTTTCCTGGGAGAACGTCACTGTGGACAAGGTTCTCGAGTT CCCATTGCTGAAAGGCCACCCTAACCTCTGTGTTCAGGTGAACAGCTCGGAGAAGCTGCA GCTGCAGGAGTGCTTGTGGGCTGACTCCCTGGGGCCTCTCAAAGACGATGTGCTACTGTT GGAGACACGAGGCCCCCAGGACAACAGATCCCTCTGTGCCTTGGAACCCAGTGGCTGTAC CCTGCAGTCAGGCCAGTGTCTGCAGCTATGGGACGATGACTTGGGAGCGCTATGGGCCTG TGCCGCTGCGCTTTCCCTCATCCTCCTTCTCAAAAAGGATCACGCGAAAGGGTGGCTGAG GCTCTTGAAACAGGACGTCCGCTCGGGGGCCGCCCAGGGGCCCGCGCGCTCTGCTCCT CTACTCAGCCGATGACTCGGGTTTCGAGCGCCTGGTGGGCGCCCTGGCGTCGGCCCTGTG CCAGCTGCCGCTGCGCGTGGCCGTAGACCTGTGGAGCCGTCGTGAACTGAGCGCGCAGGG GCCCGTGGCTTGGTTTCACGCGCAGCGGCGCCCAGACCCTGCAGGAGGGCGGCGTGGTGGT CTTGCTCTTCTCCCGGTGCGGTGGCGCTGTGCAGCGAGTGGCTACAGGATGGGGTGTC CGGGCCCGGGGCGCACGCCCGCACGACGCCTTCCGCGCCTCGCTCAGCTGCGTGCTGCC CGACTTCTTGCAGGGCCGGGCGCCCGGCAGCTACGTGGGGGCCTGCTTCGACAGGCTGCT CCACCGGACGCCGTACCCGCCCTTTTCCGCACCGTGCCCGTCTTCACACTGCCCTCCCA ACTGCCAGACTTCCTGGGGGCCCTGCAGCAGCCTCGCGCCCCGCGTTCCGGGCGGCTCCA CCCGGGGACTCCCGCGCGGGACGCGGGGTGGGACCAGGGGCGGGGACCTGGGGCGGGGA CGGGACT<u>TAA</u>ATAAAGGCAGACGCTGTTTTTCTAAAAAAA

FIGURE 364

MPVPWFLLSLALGRSPVVLSLERLVGPQDATHCSPGLSCRLWDSDILCLPGDIVPAPGPV
LAPTHLQTELVLRCQKETDCDLCLRVAVHLAVHGHWEEPEDEEKFGGAADSGVEEPRNAS
LQAQVVLSFQAYPTARCVLLEVQVPAALVQFGQSVGSVVYDCFEAALGSEVRLWSYTQPR
YEKELNHTQQLPALPWLNVSADGDNVHLVLNVSEEQHFGLSLYWNQVQGPPKPRWHKNLT
GPQIITLNHTDLVPCLCIQVWPLEPBDSVRTNICPFREDPRAHQNLWQAARLRLLTLQSWL
LDAPCSLPAEAALCWRAPGGDPCQPLVPPLSWENVTVDKVLEFPLLKGHPNLCVQVNSSE
KLQLQECLWADSLGPLKDDVLLLETRGPQDNRSLCALEPSGCTSLPSKASTRAARLGEYL
LQDLQSGQCLQLWDDDLGALWACPMDKYIHKRWALVWLACLLFAAALSLILLLKKDHAKG
WLRLLKQDVRSGAAARGRAALLLYSADDSGFERLVGALASALCQLPLRVAVDLWSREELS
AQGPVAWFHAQRQTLQEGGVVVLLFSPGAVALCSEWLQDGVSGPGAHGPHDAFRASLSC
VLPDFLQGRAPGSYVGACFDRLLHPDAVPALFRTVPVFTLPSQLPDFLGALQQPRAPRSG
RLQERAEQVSRALQPALDSYFHPPGTPAPGRGVGPGAGPGAGDGT

Signal sequence: amino acids 1-20

Transmembrane domain: amino acids 453-475

N-glycosylation sites: amino acids 118-121, 186-189, 198-201, 211-214, 238-241, 248-251, 334-337, 357-360, 391-394

Glycosaminoglycan attachment site: amino acids 583-586

cAMP- and cGMP-dependent protein kinase phosphorylation site:

N-myristoylation sites: amino acids 107-112, 152-157, 319-324, 438-443, 516-521, 612-617, 692-697, 696-701, 700-705

FIGURE 365

AATAGAAGTCCTCAGGACGGAGCAGAGGTGGCCGGCGGGCCCGGCTGACTGCCCTCTGC TTTCTTTCCATACCTTTTCTTTCGGACTCGAATCACGGCTGCTGCGAAGGGTCTAGTTC CGGACACTAGGGTGCCCGAACGCGCTGATGCCCCGAGTGCTCGCAGGGCTTCCCGCTAAC CATGCTGCCGCCGCGCGCCCGCAGCTGCCTTGGCGCTGCCTGTGCTCCTGCTACTGCT GGTGGTGCTGACGCCGCCCCGACCGGCGCAAGGCCATCCCCAGGCCCAGATTACCTGCG GCGCGGCTGGATGCGGCTGCTAGCGGAGGGCGAGGGCTGCGCTCCCTGCCGGCCAGAAGA GTGCGCCCCCCCCCGCGCCTGCCTGCCGGCAGGCTGCGCGACGCGTGCGGCTGCTGCTG GGAATGCGCCAACCTCGAGGGCCAGCTCTGCGACCTGGACCCCAGTGCTCACTTCTACGG GCACTGCGGCGAGCAGCTTGAGTGCCGGCTGGACACAGGCGGCGACCTGAGCCGCGGAGA GGTGCCGGAACCTCTGTGTGCCTGTCGTTCGCAGAGTCCGCTCTGCGGGTCCGACGGTCA CACCTACTCCCAGATCTGCCGCCTGCAGGAGGCGGCCCGCGCTCGGCCCGATGCCAACCT CACTGTGGCACACCCGGGGCCCTGCGAATCGGGGCCCCAGATCGTGTCACATCCATATGA CACTTGGAATGTGACAGGGCAGGATGTGATCTTTGGCTGTGAAGTGTTTGCCTACCCCAT GGCCTCCATCGAGTGGAGGAAGGATGGCTTGGACATCCAGCTGCCAGGGGATGACCCCCA CCAGGCTGTGCGTCCCAGTGATGAGGGCACTTACCGCTGCCTTGGCCGCAATGCCCTGGG TCAAGTGGAGGCCCCTGCTAGCTTGACAGTGCTCACACCTGACCAGCTGAACTCTACAGG TGACGATTACTAC<u>TAG</u>GTCCAGAGCTCTGGCCCATGGGGGTGGGTGAGCGGCTATAGTGT ${\tt TCATCCCTGCTCTTGAAAAGACCTGGAAAGGGGAGCAGGGTCCCTTCATCGACTGCTTTC}$ ATGCTGTCAGTAGGGATGATCATGGGAGGCCTATTTGACTCCAAGGTAGCAGTGTGGTAG GATAGAGACAAAAGCTGGAGGAGGGTAGGGAGAGAAGCTGAGACCAGGACCGGTGGGGTA CAAAGGGGCCCATGCAGGAGATGCCCTGGCCAGTAGGACCTCCAACAGGTTGTTTCCCAG GCTGGGGTGGGGCCTGAGCAGACACAGAGGTGCAGGCACCAGGATTCTCCACTTCTTCC AGCCCTGCTGGGCCACAGTTCTAACTGCCCTTCCTCCCAGGCCCTGGTTCTTGCTATTTC CTGGTCCCCAACGTTTATCTAGCTTGTTTGCCCTTTCCCCAAACTCATCTTCCAGAACTT ${\tt TTCCCTCTCTCTAAGCCCCAGTTGCACCTACTAACTGCAGTCCCTTTTGCTGTCTGCCG} \\$ TCTTTTGTACAAGAGAGAGAACAGCGGAGCATGACTTAGTTCAGTGCAGAGAGATTT

FIGURE 366

MLPPPRPAAALALPVLLLLLVVLTPPPTGARPSPGPDYLRRGWMRLLAEGEGCAPCRPEE
CAAPRGCLAGRVRDACGCCWECANLEGGLCDLDPSAHFYGHCGEQLECTDTGGDLSRGE
VPEPLCACRSQSPLCGSDGHTYSQICRLQEAARARPDANLTVAHPGPCESGFQIVSHPYD
TWNVTGQDVIFGCEVFAYPMASIEWRKDGLDIQLPGDDPHISVQFRGGPQRFEVTGWLQI
QAVRPSDEGTYRCLGRNALGQVEAPASLTVLTPDQLNSTGIFQLRSLNLVPEEEAESEEN
DDYY

Important features of the protein:

Signal peptide:

Transmembrane domain:

None

N-glycosylation site: 159-163, 183-187, 277-281

Tyrosine kinase phosphorylation site: 244-252

N-myristoylation site: 52-58, 66-72, 113-119, 249-255

Kazal-type serine protease inhibitor domain: 121-168

Immunoglobulin domain: 186-255

Insulin-like growth factor binding proteins: 53-90

FIGURE 367

FIGURE 368

MERGAGAKLLPLLLLRATGFTCAQTDGRNGYTAVIEVTSGGPWGDWAWPEMCPDGFFAS GFSLKVEPPQGIPGDDTALNGIRLHCARGNVLGNTHVVESQSGSWGEWSEPLWCRGGAYL VAFSLRVEAPTTLGDNTAANNVRFRCSDGEELQGPGLSWGDFGDWSDHCPKGACGLQTKI OGPRGLGDDTALNDARLFCCRS

Important features of the protein:

Signal peptide: 1-24

Transmembrane domain:

N-myristoylation site: 41-47 89-95 156-162

Growth factor and cytokines receptors family signature 2: 103-110

FIGURE 369

GCCAACACTGGCCAAACCTCGGAGACCGTCCTGCGCTCTCTGGAGACGCGCTGTCCGCGC CCAGGGTGGTGCCATGTGGGGGCGCTCGCCGCTCGTCCTCATCCTGGAACGCCGCGTACTGCCACGGCTGGCTGGACGCGCAGGGCGTCTGGCGCATCGGCTTCCAGTGTCCCGA GCGCTTCGACGCCGCCGCCGCCACCATCTGCTGCGCCAGCTGCGCGTTGCGCTACTGCTG CTCCAGCGCCGAGGCGCCCTGGACCAGGGCGGCTGCGACAATGACCGCCAGCAGGGCGC TGGCGAGCCTGGCCGGGCGGACAAAGACGGCCCCGACGGCTCGGCAGTGCCCATCTACGT GCCGTTCCTCATTGTTGGCTCCGTGTTTGTCGCCTTTATCATCTTGGGGTCCCTGGTGGC AGCCTGTTGCTGCAGATGTCTCCGGCCTAAGCAGGATCCCCAGCAGAGCCGAGCCCCAGG GGGTAACCGCTTGATGGAGACCATCCCCATGATCCCCAGTGCCAGCACCTCCCGGGGGTC GTCCTCACGCCAGTCCAGCACAGCTGCCAGTTCCAGCTCCAGCGCCCAACTCAGGGGCCCG GGCGCCCCAACAAGGTCACAGACCAACTGTTGCTTGCCGGAAGGGACCATGAACAACGT GTATGTCAACATGCCCACGAATTTCTCTGTGCTGAACTGTCAGCAGGCCACCCAGATTGT GCCACATCAAGGGCAGTATCTGCATCCCCCATACGTGGGGTACACGGTGCAGCACGACTC TGTGCCCATGACAGCTGTGCCACCTTTCATGGACGGCCTGCAGCCTGGCTACAGGCAGAT TCAGTCCCCCTTCCCTCACACCAACAGTGAACAGAAGATGTACCCAGCGGTGACTGTA<u>TA</u> ACCGAGAGTCACTGGTGGGTTCCTTTACTGAAGGGAGACGAAGGCAGGGGTGGATTTTCG AGGTGGAAGT

FIGURE 370

MWGARRSSVSSSWNAASLLQLLLAALLAAGARASGEYCHGWLDAQGVWRIGFQCPERFDG GDATICCGSCALRYCCSSAEARLDQGGCDNDRQQGAGEPGRADKDGFDGSAVPIYVPFLI VGSVFVAFIILGSLVAACCCRCLRPKQDPQQSRAPGGNRLMETIPMIPSASTSRGSSSRQ SSTAASSSSSANSGARAPPTRSQTNCCLPEGTMNNYVNMPTNFSVLNCQQATQIVPHQQ QYLHPPYVGYTVQHDSVPMTAVPPFMDGLQPGYRQIQSPFPHTNSEQKMYPAVTV

Important features of the protein:

Signal peptide:

1-33

Transmembrane domain:

54-78

N-qlycosylation site:

223-226

cAMP- and cGMP-dependent protein kinase phosphorylation site: 5-8

N-myristoylation site:

3-8, 30-35, 60-65, 86-91, 132-137, 211-216, 268-273

Prokaryotic membrane lipoprotein lipid attachment site: 128-138

FIGURE 371

CACCAGACAGCACTCCAGCACTCTGTTTGGGGGGCATTCGAAACAGCAAAATCACTCATA AAAGGCAAAAAATTGCAAAAAAAAATAGTAATAACCAGCATGGCACTAAATAGACCATGA AAAGACATGTGTGTGCAGTATGAAAATTGAGACAGGAAGGCAGAGTGTCAGCTTGTTCCA CCTCAGCTGGGA<u>ATC</u>TGCATCAGGCAACTCAAGTTTTTCACCACGGCATGTGTCTGTGAA TGTCCGCAAAACATTCTCTCTCCCCAGCCTTCATGTGTTAACCTGGGGATGATGTGGACC TGGGCACTGTGGATGCTCCCTTCACTCTGCAAATTCAGCCTGGCAGCTCTGCCAGCTAAG CCTGAGAACATTTCCTGTGTCTACTACTATAGGAAAAATTTAACCTGCACTTGGAGTCCA GGAAAGGAAACCAGTTATACCCAGTACACAGTTAAGAGAACTTACGCTTTTGGAGAAAAA CATGATAATTGTACAACCAATAGTTCTACAAGTGAAAATCGTGCTTCGTGCTCTTTTTTC CTTCCAAGAATAACGATCCCAGATAATTATACCATTGAGGTGGAAGCTGAAAATGGAGAT GGTGTAATTAAATCTCATATGACATACTGGAGATTAGAGAACATAGCGAAAACTGAACCA CCTAAGATTTTCCGTGTGAAACCAGTTTTGGGCATCAAACGAATGATTCAAATTGAATGG ATAAAGCCTGAGTTGGCGCCTGTTTCATCTGATTTAAAATACACACTTCGATTCAGGACA GTCAACAGTACCAGCTGGATGGAAGTCAACTTCGCTAAGAACCGTAAGGATAAAAACCAA ACGTACAACCTCACGGGGCTGCAGCCTTTTACAGAATATGTCATAGCTCTGCGATGTGCG GTCAAGGAGTCAAAGTTCTGGAGTGACTGGAGCCAAGAAAAAATGGGAATGACTGAGGAA GAAGCTCCATGTGGCCTGGAACTGTGGAGAGTCCTGAAACCAGCTGAGGCGGATGGAAGA AGGCCAGTGCGGTTGTTATGGAAGAAGGCAAGAGGGCCCCAGTCCTAGAGAAAACACTT GGCTACAACATATGGTACTATCCAGAAAGCAACACTAACCTCACAGAAACAATGAACACT ACTAACCAGCAGCTTGAACTGCATCTGGGAGGCGAGAGCTTTTGGGTGTCTATGATTTCT TATAATTCTCTTGGGAAGTCTCCAGTGGCCACCCTGAGGATTCCAGCTATTCAAGAAAAA TCATTTCAGTGCATTGAGGTCATGCAGGCCTGCGTTGCTGAGGACCAGCTAGTGGTGAAG TGGCAAAGCTCTGCTCTAGACGTGAACACTTGGATGATTGAATGGTTTCCGGATGTGGAC TCAGAGCCCACCACCCTTTCCTGGGAATCTGTGTCTCAGGCCACGAACTGGACGATCCAG CAAGATAAATTAAAACCTTTCTGGTGCTATAACATCTCTGTGTATCCAATGTTGCATGAC AAAGTTGGCGAGCCATATTCCATCCAGGCTTATGCCAAAGAAGGCGTTCCATCAGAAGGT CCTGAGACCAAGGTGGAGAACATTGGCGTGAAGACGGTCACGATCACATGGAAAGAGATT CCCAAGAGTGAGAAAAGGGTATCATCTGCAACTACACCATCTTTTACCAAGCTGAAGGT GGAAAAGGATTCTGTAAGCACGCCCATAGCGAAGTGGAAAAAAACCCCAAGCCCCAGATA GATGCTATGGATAGACCTGTTGTAGGCATGGCTCCCCCATCTCATTGTGACTTGCAACCT TTGGGGTTTTGGGGGTTAAATGAGAGTGAAGTGACAGTACCTGAGAGGAGAGTCCTGAGG AAATGGAAGGAGTTGTTA<u>TAA</u>TTTGTCCTGGTTAGGCCCTGAATTGACCTCCCGGGAGCT CCCCGACCATCATTCCCAGGAATGGCGTGCCTGGCTTAAAGAGTGAGGAGGAACAGACCC TGTCACCATGACTTCTACTGCCCCTGCCAAATCATGCTTTTGTTTTTCAGTCCACCTTAT CTCCTGACATCTTAAATACTGGGCAAGGCTTGGATTCTTGCTTAGGCTAAATAATTTTTT CTTATGGTAAAATACACGTAAAATATTTTTCCAGTTTAAACATTTGAAAGTGTACAATTT AGTGGCATTAGAAGCATTCACAATATTGTGCAACCATCACCACTATTTCCAGAACTCTTC TATTTCTGCCCAAATAGAAGCCCTATACCCATTCATTAGTCACTCCCCATTCCTCCTC CCACAGCCCCTGGCAACTACCAAACTGCTTTGTGTCTCTATGGATTGCCTATTTTGGATA

FIGURE 372

MCIRQLKFFTTACVCECPQNILSPQPSCVNLGMMWTWALWMLPSLCKFSLAALPAKPENI
SCVYYYRKNLTCTWSPGKETSYTQYTVKRTYAFGEKHDNCTTNSSTSENRASCSFFLOFI
TIPDNYTIEVEAENGDGVIKSHMTYWRLENIAKTEPPKIFRVKPVLGIKRMIQIEWIKPE
LAPVSSDLKYTLRFRTVNSTSWMEVNFAKNRKDKNQTYNLTGLQPFTEYVIALRCAVKES
KFWSDWSQEKMGMTEEEAPCGLELWRVLKPAEADGRPVRLLWKKARGAPVLEKTLGYNI
WYYPESNTNLTETMNTTNQLELHLGGESFWVSMISYNSLGKSPVATLRIPAIQEKSFQC
IEVMQACVAEDQLVVKWQSSALDVNTWMIEWFPDVDSEPTTLSWESVSQATNWTIQODKL
KPFWCYNISVYPMLHDKVGEPYSIQAYAKEGVPSEGPETKVENIGVKTVTTTWKEIPKSE
RKGIICNTTIFYQAEGGKGFCKHAHSEVEKNPKPQIDAMDRPVVGMAPPSHCDLQPGMNH
LASINLSRMGAKSTHLLGFWGLBESEVTVPERRVLRKWKELL

Important features of the protein:

Signal peptide:

1-46

Transmembrane domain:

None

N-glycosylation site:

59-63, 69-73, 99-103, 103-107, 125-129, 198-202, 215-219, 219-223, 309-313, 315-319, 412-416, 427-431, 487-491, 545-549, 563-567

N-myristoylation site:

32-38, 137-143, 483-489, 550-556, 561-567

Amidation site:

274-278

Growth factor and cytokines receptors family signature 1: 62-75

Fibronectin type III domain:

54-144

154-247

FIGURE 373

CCAGGTCCAACTGCACCTCGGTTCTATCGATTGAATTCCCCGGGGATCCTCTAGAGATCC CTCGACCTCGACCCACGCGTCCGCCAAGCTGGCCCTGCACGGCTGCAAGGGAGGCTCCTG TGGACAGGCCAGGCAGGTGGGCCTCAGGAGGTGCCTCCAGGCGGCCAGTGGGCCTGAGGC GCCTGGGCTCCAGCAGCATCAGCAGCCCCCAGGACCGGGGGAGGCACAGGTGGCCCCCAC ACCCAGAGGAGAAGGCCACCCGCCTGGAGGCACAGGCCATGAGGGGCTCTCAGGAGGTG CTGCTGATGTGGCTTCTGGTGTTGGCAGTGGGCGGCACAGAGCACGCCTACCGGCCCGGC GTGTGTACCAGCCTTCCTCACCACCTGCGACGGGCACCGGGCCTGCAGCACCTACCGAA CCATTTATAGGACCGCCTACCGCCGCAGCCCTGGGCTGGCCCCTGCCAGGCCTCGCTACG CGTGCTGCCCGGCTGGAAGAGGACCAGCGGGCTTCCTGGGGCCTGTGGAGCAGCAATAT GCCAGCCGCATGCCGGAACGGAGGGAGCTGTCCAGCCTGGCCGCTGCCGCTGCCCTG CAGGATGGCGGGGTGACACTTGCCAGTCAGATGTGGATGAATGCAGTGCTAGGAGGGGCG GCTGTCCCCAGCGCTGCATCAACACCGCCGGCAGTTACTGGTGCCAGTGTTGGGAGGGGC CCAACCCGACAGGAGTGGACAGTGCAATGAAGGAAGAAGTGCAGAGGCTGCAGTCCAGGG TGGACCTGCTGGAGGAGAAGCTGCAGCTGGTGCTGGCCCCACTGCACAGCCTGGCCTCGC AGGCACTGGAGCATGGGCTCCCGGACCCCGGCAGCCTCCTGGTGCACTCCTTCCAGCAGC GCTCCTGCAAGAAAGACTCGTGACTGCCCAGCGCCCCAGGCTGGACTGAGCCCCTCACGC CGCCCTGCAGCCCCATGCCCCTGCCCAACATGCTGGGGGTCCAGAAGCCACCTCGGGGT GACTGAGCGGAAGGCCAGGCAGGCCTTCCTCCTTTTCCTCCTCCCCTTCCCTCGGGAGG GTCCCAGACCCTGGCATGGGATGGGCTGGGATTTTTTTTGTGAATCCACCCCTGGCTAC CCCCACCCTGGTTACCCCAACGGCATCCCAAGGCCAGGTGGGCCCTCAGCTGAGGGAAGG TACGAGTTCCCCTGCTGGAGCCTGGGACCCATGGCACAGGCCAGGCAGCCCGGAGGCTGG GTGGGGCCTCAGTGGGGGCTGCTGCCTGACCCCCAGCACAATAAAAATGAAACGTGAAAA GACCTGCAGAAGCTTGGCCGCCATGGCCCAACTTGTTTATTGCAGCTTATAATGGTTACAAAT

FIGURE 374

MTDSPPPGHPEEKATPPGGTGHEGLSGGAADVASGVGSGRHRARLPARPLGCVLSRAHGD PVSESFVQRVYQPFLTTCDGHRACSTYRTIYRTAYRRSPGLAPARPRYACCPGWKRTSGL PGACGAAICQPPCRNGGSCVQPGRCRCPAGWRGDTCQSDVDECSARRGGCPQRCINTAGS YWCQCWEGHSLSADGTLCVPKGGPPRVAPNPTGVDSAWKEEVQRLQSRVDLLEEKLQLVL APLHSLASQALEHGLPPPGSLLVHSFQQLGRIDSLSEQISFLEEQLGSCSCKKDS

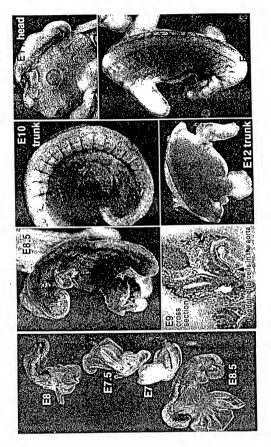
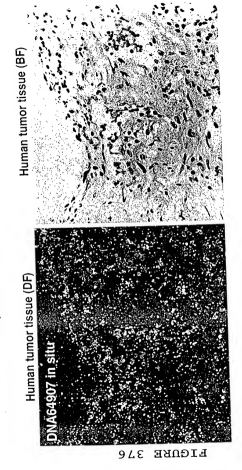


FIGURE 375



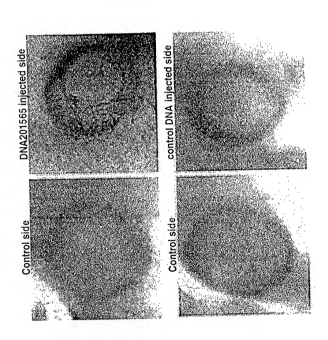


FIGURE 377